2	UTILITY PATENT APPLICATION TRANSMITTAL			
į	Submit an original and a duplicate for fee processing (Only for new nonprovisional applications under 37 CFR 1.53(b))			
3	ADDRESS TO:	Attorney Docket No. MBHB00-1213		
		First Named Inventor Saris et al		
	Assistant Commissioner for Patents Box Patent Application Washington, D.C. 20231	Express Mail No. EL625499230US		
	7740g.c., 510. 2020.	Total Pages		
į	APPLICATION ELEMENTS	ACCOMPANYING APPLICATION PARTS		
	 Transmittal Form with Fee Specification (including claims and 	Assignment Papers Power of Attorney		
	abstract) [Total Pages 5] 3. Drawings [Total Sheets 27]	10. English Translation Document (if applicable) 11. Information Disclosure Statement (IDS)		
	4. Oath or Declaration Total Pages]	☐ PTO-1449 Form		
	a. Newly executedb. Copy from prior application	Copies of IDS Citations 12. Preliminary Amendment		
	[Note Boxes 5 and 17 below]	13. ⊠ Return Receipt Postcard		
	i. Deletion of Inventor(s) Signed	(Should be specifically itemized) 14. Small Entity Statement(s)		
	statement attached deleting inventor(s) named in the prior application	Enclosed		
	5. Incorporation by Reference: The entire disclosure of the prior application, from which a	Statement filed in prior application; status still proper and desired		
	copy of the oath or declaration is supplied under Box 4b, is considered as being part of the	15. Certified Copy of Priority Document(s)		
	disclosure of the accompanying application and is hereby incorporated by reference therein.	16. 🛛 Other: Title Page		
	6. Microfiche Computer Program			
	 Nucleotide and/or Amino Acid Sequence Submission 			
	a. 🛛 Computer Readable Copy			
	b. ⊠ Paper Copyc. ⊠ Statement verifying above copies			
17. A This is a CONTINUING APPLICATION. Please note the following:				
	a. ☑ This is a ☑ Continuation ☐ Divisional ☐ Continuation-in-part			
	of prior application Serial No. 60/170,191 b. Cancel in this application original claimsof the prior application before calculating the filing fee. c. Amend the specification by inserting before the first line the sentence: This is a continuation divisional continuation-in-part of application Serial No.			
1	d The prior application is assigned or	frecord to Amgen Inc		



Marie de la companya de la companya

UTILITY PATEN	UTILITY PATENT APPLICATION TRANSMITTAL Attorney Docket No. MBHB00-1215					
APPLICATION FEES						
BASIC FEE				710,00		
CLAIMS	3 37 37 38 38 2 38 32	NUMBER EXTRA	RATE			
Total Claims	112-20=	92	x \$18.00 \$.00		
Independent Clair	ms 10-3=	7	x \$80.00 \$	560.00		
	ndent Claims(s) if applicable		+\$270.00 \$	270.00		
		Total of abov		.00		
1504000 ASS ASS ASS ASS ASS ASS ASS ASS ASS		eduction by 50% for filing t		1/2		
Assignment fe	ee if applicable		+ \$40.00 \$.00		
			TOTAL = \$	3,196.00		
	charge my Deposit Account N	No. 13-2490 in the amount	of \$			
19. 🛛 A chec	k in the amount of \$ 3,196.00	is enclosed.				
20. The Commiss	sioner is hereby authorized to	credit overpayments or ch	harge any additional	fees of the		
	es to Deposit Account No. 13-		0 ,			
	ees required under 37 CFR 1					
	ees required under 37 CFR 1					
c. 🖾 F	ees required under 37 CFR 1	.18.		•		
21. The Commissioner is hereby generally authorized under 37 CFR 1.136(a)(3) to treat any future reply in this or any related application filed pursuant to 37 CFR 1.53 requiring an extension of time as incorporating a request therefor, and the Commissioner is hereby specifically authorized to charge Deposit Account No. 13-2490 for any fee that may be due in connection with such a request for an extension of time.						
<u> </u>	22. CERT	IFICATE OF MAILING				
I hereby certify that I directed that the correspondence identified above be deposited with the United States Postal Service as "Express Mail Post Office to Addressee" under 37 CFR § 1.10 on the date indicated below and is addressed to the Asst. Commissioner for Patents, Box Patent Application, Washington, DC 20231.						
23. USPTO CUSTOMER NUMBER						
PATENT & TRADEHARK OFFICE O20306						
24. CORRESPONDENCE ADDRESS						
OF CIONATURE OF ARRUGANT ATTORNEY OR ACENT REQUIRED						
25. SIGNATURE OF APPLICANT, ATTORNEY, OR AGENT REQUIRED						
Name Reg. No.	Kevin E Noonan, Reg. No.3	5,303				
Signature /						
6	//	·				
Date	November 28, 2000					

APPLICATION FOR UNITED STATES LETTERS PATENT IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

(Case No. 00,1213)

Inventors:

Christiaan M. Saris 4027 Colonett Place

Newbury Park, CA 91320

Citizen of the Netherlands

Jennifer Giles

3031 Charlotte Street

Newbury Park, CA 91320

Citizen of the United States of America

Sharon X. Mu 2954 Rikkard Drive

Thousand Oaks, CA 91362

Citizen of the United States of America

Min Xia

3946 Calle Buena Vista Newbury Park, CA 91320

Citizen of Canada

Michael Brian Bass 1743 N. Marian Avenue Thousand Oaks, CA 91360

Citizen of the United States of America

Roger Craveiro 384 East Wilbur Road, Apt. 203 Thousand Oaks, CA 91360

Citizen of the United States of America

Assignee:

Amgen, Inc.

A Corporation of the State of Delaware

Title:

Interleukin-1 Receptor Antagonist-Related Molecules and Uses Thereof

INTERLEUKIN-1 RECEPTOR ANTAGONIST-RELATED MOLECULES AND USES THEREOF

Field of the Invention

5

10

15

20

25

30

The present invention relates to novel Interleukin-1 Receptor Antagonist-Related (IL-1ra-R) polypeptides and nucleic acid molecules encoding the same. The invention also relates to selective binding agents, vectors, host cells, and methods for producing IL-1ra-R polypeptides. The invention further relates to pharmaceutical compositions and methods for the diagnosis, treatment, amelioration, and/or prevention of diseases, disorders, and conditions associated with IL-1ra-R polypeptides.

Background of the Invention

Technical advances in the identification, cloning, expression, and manipulation of nucleic acid molecules and the deciphering of the human genome have greatly accelerated the discovery of novel therapeutics. Rapid nucleic acid sequencing techniques can now generate sequence information at unprecedented rates and, coupled with computational analyses, allow the assembly of overlapping sequences into partial and entire genomes and the identification of polypeptide-encoding regions. A comparison of a predicted amino acid sequence against a database compilation of known amino acid sequences allows one to determine the extent of homology to previously identified sequences and/or structural landmarks. The cloning and expression of a polypeptide-encoding region of a nucleic acid molecule provides a polypeptide product for structural and functional analyses. The manipulation of nucleic acid molecules and encoded polypeptides may confer advantageous properties on a product for use as a therapeutic.

In spite of the significant technical advances in genome research over the past decade, the potential for the development of novel therapeutics based on the human genome is still largely unrealized. Many genes encoding potentially beneficial polypeptide therapeutics or those encoding polypeptides, which may

act as "targets" for therapeutic molecules, have still not been identified.

Accordingly, it is an object of the invention to identify novel polypeptides, and nucleic acid molecules encoding the same, which have diagnostic or therapeutic benefit.

5

10

15

20

25

30

One of the most potent inflammatory cytokines yet discovered is interleukin-1 (IL-1). IL-1 is thought to be involved in many diseases and medical conditions. It is produced (though not exclusively) by cells of the macrophage/monocyte lineage, and may be produced in two forms: IL-1alpha (IL-1 Δ) and IL-1beta (IL-1E). Interleukin-1 receptor antagonist (IL-1ra) is a human protein that acts as a natural inhibitor of interleukin-1.

Summary of the Invention

The present invention relates to novel IL-1ra-R nucleic acid molecules and encoded polypeptides.

The invention provides for an isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:

- (a) the nucleotide sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35;
- (b) the nucleotide sequence of the DNA insert in ATCC Deposit No. PTA-1423;
- (c) a nucleotide sequence encoding the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (d) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (c); and
 - (e) a nucleotide sequence complementary to any of (a) (c).

The invention also provides for an isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:

(a) a nucleotide sequence encoding a polypeptide which is at least about 70 percent identical to the polypeptide as set forth in any of SEQ ID NO: 2,

10

15

20

25

SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;

- (b) a nucleotide sequence encoding an allelic variant or splice variant of the nucleotide sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the nucleotide sequence of the DNA insert in ATCC Deposit No. PTA-1423, or (a);
- (c) a region of the nucleotide sequence of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the DNA insert in ATCC Deposit No. PTA-1423, (a), or (b) encoding a polypeptide fragment of at least about 25 amino acid residues, wherein the polypeptide fragment has an activity of the encoded polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or is antigenic;
- (d) a region of the nucleotide sequence of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the DNA insert in ATCC Deposit No. PTA-1423, or any of (a) (c) comprising a fragment of at least about 16 nucleotides;
- (e) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (d); and
 - (f) a nucleotide sequence complementary to any of (a) (d).

The invention further provides for an isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:

- (a) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one conservative amino acid substitution, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (b) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid insertion, wherein the encoded polypeptide has an activity of the

polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;

- (c) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid deletion, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (d) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 which has a C-and/or N- terminal truncation, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (e) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one modification selected from the group consisting of amino acid substitutions, amino acid insertions, amino acid deletions, C-terminal truncation, and N-terminal truncation, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (f) a nucleotide sequence of any of (a) (e) comprising a fragment of at least about 16 nucleotides;
- (g) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (f); and
 - (h) a nucleotide sequence complementary to any of (a) (e).

25

30

5

10

15

20

The present invention provides for an isolated polypeptide comprising an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36; and
- (b) the amino acid sequence encoded by the DNA insert in ATCC Deposit No. PTA-1423.

The invention also provides for an isolated polypeptide comprising the amino acid sequence selected from the group consisting of:

- (a) an amino acid sequence for an ortholog of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (b) an amino acid sequence which is at least about 70 percent identical to the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (c) a fragment of the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 comprising at least about 25 amino acid residues, wherein the fragment has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or is antigenic; and
- (d) an amino acid sequence for an allelic variant or splice variant of the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, the amino acid sequence encoded by the DNA insert in ATCC Deposit No. PTA-1423, (a), or (b).

20

25

30

15

5

10

The invention further provides for an isolated polypeptide comprising the amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one conservative amino acid substitution, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (b) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid insertion, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;

- (c) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid deletion, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (d) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 which has a C- and/or N- terminal truncation, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36; and
- (e) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one modification selected from the group consisting of amino acid substitutions, amino acid insertions, amino acid deletions, C-terminal truncation, and N-terminal truncation, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

20

25

30

10

5

The invention still further provides for an isolated polypeptide comprising the amino acid sequence as set forth in either SEQ ID NO: 1 or SEQ ID NO: 3 with at least one amino acid substitution selected from the group consisting of: arginine at position 2; alanine, lysine, or arginine at position 3; serine at position 7; lysine at position 8; alanine, cysteine, lysine, threonine, or serine at position 9; cysteine or phenylalanine at position 10; arginine or trptophan at position 13; serine at position 15; arginine at position 18; serine or threonine at position 19; threonine at position 21; serine at position 23; arginine at position 34; tyrosine, serine, or arginine at position 37; lysine, arginine, threonine, or serine at position 38; threonine at position 41; serine, phenylalanine, or alanine at position 43; alanine at position 44; serine or lysine at position 48; alanine, threonine, or phenylalanine at position 52; serine at position 53; serine at position 54; alanine or tyrosine at position 58; lysine at position 65; phenylalanine at position 66; tyrosine at position 67; serine, tyrosine, or phenylalanine at position 69; lysine or serine at position 73; threonine or arginine at position 78; serine or alanine at position 90; alanine at position 91; serine at position 96; lysine or arginine at

10

15

20

25

30

position 97; lysine or serine at position 98; alanine at position 100; tyrosine at position 102; arginine or alanine at position 104; lysine at position 105; threonine at position 106; arginine at position 108; lysine, threonine, or trptophan at position 109; threonine or serine at position 110; serine at position 111; serine at position 114; serine at position 116; phenylalanine, cysteine, or tyrosine at position 117; tyrosine at position 121; serine or alanine at position 123; cysteine, serine, or threonine at position 126; serine at position 136; phenylalanine or arginine at position 138; threonine, arginine, or alanine at position 141; lysine or tyrosine at position 142; trptophan or threonine at position 143; alanine at position 145; threonine or serine at position 147; cysteine at position 151; and serine, cysteine, or phenylalanine at position 152; wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, or SEQ ID NO: 6.

The invention still further provides for an isolated polypeptide comprising the amino acid sequence as set forth in SEQ ID NO: 5 with at least one amino acid substitution selected from the group consisting of: arginine at position 21; alanine, lysine, or arginine at position 22; serine at position 26; lysine at position 27; alanine, cysteine, lysine, threonine, or serine at position 28; cysteine or phenylalanine at position 29; arginine or trptophan at position 32; serine at position 34; arginine at position 37; serine or threonine at position 38; threonine at position 40; serine at position 42; arginine at position 53; tyrosine, serine, or arginine at position 56; lysine, arginine, threonine, or serine at position 57; threonine at position 60; serine, phenylalanine, or alanine at position 62; alanine at position 63; serine or lysine at position 67; alanine, threonine, or phenylalanine at position 71; serine at position 72; serine at position 73; alanine or tyrosine at position 77; lysine at position 84; phenylalanine at position 85; tyrosine at position 86; serine, tyrosine, or phenylalanine at position 88; lysine or serine at position 92; threonine or arginine at position 97; serine or alanine at position 109; alanine at position 110; serine at position 115; lysine or arginine at position 116; lysine or serine at position 117; alanine at position 119; tyrosine at position 121; arginine or alanine at position 123; lysine at position 124; threonine at position

10

15

20

25

125; arginine at position 127; lysine, threonine, or trptophan at position 128; threonine or serine at position 129; serine at position 130; serine at position 133; serine at position 135; phenylalanine, cysteine, or tyrosine at position 136; tyrosine at position 140; serine or alanine at position 142; cysteine, serine, or threonine at position 145; serine at position 155; phenylalanine or arginine at position 157; threonine, arginine, or alanine at position 160; lysine or tyrosine at position 161; trptophan or threonine at position 162; alanine at position 164; threonine or serine at position 166; cysteine at position 170; and serine, cysteine, or phenylalanine at position 171; wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, or SEQ ID NO: 6.

Also provided are fusion polypeptides comprising IL-1ra-R amino acid sequences.

The present invention also provides for an expression vector comprising the isolated nucleic acid molecules as set forth herein, recombinant host cells comprising the recombinant nucleic acid molecules as set forth herein, and a method of producing an IL-1ra-R polypeptide comprising culturing the host cells and optionally isolating the polypeptide so produced.

A transgenic non-human animal comprising a nucleic acid molecule encoding an IL-1ra-R polypeptide is also encompassed by the invention. The IL-1ra-R nucleic acid molecules are introduced into the animal in a manner that allows expression and increased levels of an IL-1ra-R polypeptide, which may include increased circulating levels. Alternatively, the IL-1ra-R nucleic acid molecules are introduced into the animal in a manner that prevents expression of endogenous IL-1ra-R polypeptide (*i.e.*, generates a transgenic animal possessing an IL-1ra-R polypeptide gene knockout). The transgenic non-human animal is preferably a mammal, and more preferably a rodent, such as a rat or a mouse.

Also provided are derivatives of the IL-1ra-R polypeptides of the present invention.

The state of the s

5

10

15

20

25

30

Additionally provided are selective binding agents such as antibodies and peptides capable of specifically binding the IL-1ra-R polypeptides of the invention. Such antibodies and peptides may be agonistic or antagonistic.

Pharmaceutical compositions comprising the nucleotides, polypeptides, or selective binding agents of the invention and one or more pharmaceutically acceptable formulation agents are also encompassed by the invention. The pharmaceutical compositions are used to provide therapeutically effective amounts of the nucleotides or polypeptides of the present invention. The invention is also directed to methods of using the polypeptides, nucleic acid molecules, and selective binding agents.

The IL-1ra-R polypeptides and nucleic acid molecules of the present invention may be used to treat, prevent, ameliorate, and/or detect diseases and disorders, including those recited herein.

The present invention also provides a method of assaying test molecules to identify a test molecule that binds to an IL-1ra-R polypeptide. The method comprises contacting an IL-1ra-R polypeptide with a test molecule to determine the extent of binding of the test molecule to the polypeptide. The method further comprises determining whether such test molecules are agonists or antagonists of an IL-1ra-R polypeptide. The present invention further provides a method of testing the impact of molecules on the expression of IL-1ra-R polypeptide or on the activity of IL-1ra-R polypeptide.

Methods of regulating expression and modulating (i.e., increasing or decreasing) levels of an IL-1ra-R polypeptide are also encompassed by the invention. One method comprises administering to an animal a nucleic acid molecule encoding an IL-1ra-R polypeptide. In another method, a nucleic acid molecule comprising elements that regulate or modulate the expression of an IL-1ra-R polypeptide may be administered. Examples of these methods include gene therapy, cell therapy, and anti-sense therapy as further described herein.

In another aspect of the present invention, the IL-1ra-R polypeptides may be used for identifying receptors thereof ("IL-1ra-R polypeptide receptors"). Various forms of "expression cloning" have been extensively used to clone

receptors for protein ligands. See, e.g., Simonsen and Lodish, 1994, Trends Pharmacol. Sci. 15:437-41 and Tartaglia et al., 1995, Cell 83:1263-71. The isolation of an IL-1ra-R polypeptide receptor is useful for identifying or developing novel agonists and antagonists of the IL-1ra-R polypeptide signaling pathway. Such agonists and antagonists include soluble IL-1ra-R polypeptide receptors, anti-IL-1ra-R polypeptide receptor-selective binding agents (such as antibodies and derivatives thereof), small molecules, and antisense oligonucleotides, any of which can be used for treating one or more disease or disorder, including those disclosed herein.

10

5

Brief Description of the Figures

Figures 1A-1B illustrate the nucleotide sequence of the human IL-1ra-R gene (SEQ ID NO: 1) and the deduced amino acid sequence of human IL-1ra-R polypeptide (SEQ ID NO: 2);

15

Figures 2A-2B illustrate the nucleotide sequence of a human IL-1ra-R gene variant (SEQ ID NO: 3) and the deduced amino acid sequence of the human IL-1ra-R polypeptide (SEQ ID NO: 4) encoded by this vaiant;

Figure 3 illustrates the nucleotide sequence of a human IL-1ra-R splice variant (SEQ ID NO: 5) and the deduced amino acid sequence of the human IL-1ra-R polypeptide (SEQ ID NO: 6) encoded by this splice variant;

Figures 4A-4B illustrate the amino acid sequence alignment of human IL-1Δ (IL-1_alpha; SEQ ID NO: 7), human IL-1E (IL-1_beta; SEQ ID NO: 8), human IL-1 receptor antagonist (IL-1RA; SEQ ID NO: 9), human IL-1Γ (IL-1_delta; SEQ ID NO: 10), human IL-1ra-R polypeptide (IL-1ra-R; SEQ ID NO: 2), human Tango-77 (Tango-77; SEQ ID NO: 11), human Zilla4 (Zilla4; SEQ ID NO: 12), human IL-1] (IL-1_zeta; SEQ ID NO: 13), human IL-1 receptor antagonist E (IL-1 alpha; SEQ ID NO: 14), human SPOIL II (Spoil_II; SEQ ID NO: 15), human

15

IL-1H(IL-1_epsilon; SEQ ID NO: 16), and human IL-1K (IL-1_eta; SEQ ID NO: 17);

Figure 5 schematically illustrates the phylogenetic relationship of the IL-1ra gene family.

Figure 6 schematically illustrates the relationship between human IL-1ra-R polypeptide (Mature CS329), the sequence variant of human IL-1ra-R polypeptide (Mature CS329 Variant protein), and the splice variant of human IL-1ra-R polypeptide (Omega 329 protein);

Figure 7 illustrates the nucleotide sequence of the murine IL-1ra-R gene (SEQ ID NO: 35) and the deduced amino acid sequence of murine IL-1ra-R polypeptide (SEQ ID NO: 36);

Figure 8 illustrates the amino acid sequence alignment of human IL-1ra-R polypeptide (huIL-1ra-R; SEQ ID NO: 2) and murine IL-1ra-R polypeptide (muIL-1ra-R; SEQ ID NO: 36);

Figures 9A-9I illustrate the genomic nucleotide sequence for the murine IL-1ra-R gene (SEQ ID NO: 37). The locations of the coding portions of exons 1-4 are indicated (underline);

Figures 10A-10C illustrate the expression of murine (Figure 10A) and human (Figures 10B-10C) IL-1ra-R mRNA as detected by Northern blot analysis;

Figures 11A-11B illustrate the results of Western blot analysis using an anti-IL-1ra-R antibody;

Figure 12 illustrates the results of FACS analysis of spleen and bone marrow cells recovered from lethally irradiated recipient mice transplanted with murine bone

marrow cells transduced with a retroviral vector containing the IL-1ra-R gene. The percentage of cells within the lymphoid compartment is shown for cells subjected to standard FACS analysis using the indicated cell surface markers;

Figures 13A-13B illustrate the results of colony assays performed on spleen and bone marrow cells recovered from lethally irradiated recipient mice transplanted with murine bone marrow cells transduced with a retroviral vector containing the IL-1ra-R gene. Bone marrow and spleen cells from transduced and control mice were cultured under standard colony assay conditions and colonies were counted on day 14;

Figure 14 illustrates the production of ϑ -interferon in response to IL-12 treatment in spleen cells recovered from lethally irradiated recipient mice transplanted with murine bone marrow cells transduced with a retroviral vector containing the IL-1ra-R gene. A sample of conditioned media was removed from cultures (2 x 10^6 cells/well/ml) grown in the presence of IL-12 (1 ng/ml) for 48 hours and IFN- ϑ was quantitated by ELISA;

Figure 15 illustrates the production of 9-interferon in response to IL-12 and IL-18 treatment in spleen cells recovered from lethally irradiated recipient mice transplanted with murine bone marrow cells transduced with a retroviral vector containing the IL-1ra-R gene. A sample of conditioned media was removed from cultures (2 x 10⁶ cells/well/ml) grown in the presence of IL-12 (1 ng/ml) and IL-18 (10 ng/ml) for 48 hours and IFN-9 was quantitated by ELISA.

25

20

15

Detailed Description of the Invention

The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described. All references cited in this application are expressly incorporated by reference herein.

30

Definitions

15

20

25

30

The terms "IL-1ra-R gene" or "IL-1ra-R nucleic acid molecule" or "IL-1ra-R polynucleotide" refer to a nucleic acid molecule comprising or consisting of a nucleotide sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, a nucleotide sequence encoding the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, a nucleotide sequence of the DNA insert in ATCC Deposit No. PTA-1423, and nucleic acid molecules as defined herein.

The term "IL-1ra-R polypeptide allelic variant" refers to one of several possible naturally occurring alternate forms of a gene occupying a given locus on a chromosome of an organism or a population of organisms.

The term "IL-1ra-R polypeptide splice variant" refers to a nucleic acid molecule, usually RNA, which is generated by alternative processing of intron sequences in an RNA transcript of IL-1ra-R polypeptide amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

The term "isolated nucleic acid molecule" refers to a nucleic acid molecule of the invention that (1) has been separated from at least about 50 percent of proteins, lipids, carbohydrates, or other materials with which it is naturally found when total nucleic acid is isolated from the source cells, (2) is not linked to all or a portion of a polynucleotide to which the "isolated nucleic acid molecule" is linked in nature, (3) is operably linked to a polynucleotide which it is not linked to in nature, or (4) does not occur in nature as part of a larger polynucleotide sequence. Preferably, the isolated nucleic acid molecule of the present invention is substantially free from any other contaminating nucleic acid molecule(s) or other contaminants that are found in its natural environment that would interfere with its use in polypeptide production or its therapeutic, diagnostic, prophylactic or research use.

The term "nucleic acid sequence" or "nucleic acid molecule" refers to a DNA or RNA sequence. The term encompasses molecules formed from any of the known base analogs of DNA and RNA such as, but not limited to 4-acetylcytosine, 8-hydroxy-N6-methyladenosine, aziridinyl-cytosine,

10

15

20

25

3.0

uracil. 5-fluorouracil, 5-5-(carboxyhydroxylmethyl) pseudoisocytosine, 5-carboxymethylaminomethyl-2-thiouracil, 5-carboxybromouracil. methylaminomethyluracil, dihydrouracil, inosine, N6-iso-pentenyladenine, 1methyladenine, 1-methylpseudouracil, 1-methylguanine, 1-methylinosine, 2,2dimethyl-guanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-7-methylguanine, N6-methyladenine, methylcytosine, beta-D-5-methoxyamino-methyl-2-thiouracil, methylaminomethyluracil, mannosylqueosine, 5' -methoxycarbonyl-methyluracil, 5-methoxyuracil, 2methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid methylester, uracil-5oxyacetic acid, oxybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, N-uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid, pseudouracil, queosine, 2-thiocytosine, and 2.6-diaminopurine.

The term "vector" is used to refer to any molecule (e.g., nucleic acid, plasmid, or virus) used to transfer coding information to a host cell.

The term "expression vector" refers to a vector that is suitable for transformation of a host cell and contains nucleic acid sequences that direct and/or control the expression of inserted heterologous nucleic acid sequences. Expression includes, but is not limited to, processes such as transcription, translation, and RNA splicing, if introns are present.

The term "operably linked" is used herein to refer to an arrangement of flanking sequences wherein the flanking sequences so described are configured or assembled so as to perform their usual function. Thus, a flanking sequence operably linked to a coding sequence may be capable of effecting the replication, transcription and/or translation of the coding sequence. For example, a coding sequence is operably linked to a promoter when the promoter is capable of directing transcription of that coding sequence. A flanking sequence need not be contiguous with the coding sequence, so long as it functions correctly. Thus, for example, intervening untranslated yet transcribed sequences can be present between a promoter sequence and the coding sequence and the promoter sequence can still be considered "operably linked" to the coding sequence.

10

15

20

25

30

The term "host cell" is used to refer to a cell which has been transformed, or is capable of being transformed with a nucleic acid sequence and then of expressing a selected gene of interest. The term includes the progeny of the parent cell, whether or not the progeny is identical in morphology or in genetic make-up to the original parent, so long as the selected gene is present.

The term "IL-1ra-R polypeptide" refers to a polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 and related polypeptides. Related polypeptides include IL-1ra-R polypeptide fragments, IL-1ra-R polypeptide orthologs, IL-1ra-R polypeptide variants, and IL-1ra-R polypeptide derivatives, which possess at least one activity of the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. IL-1ra-R polypeptides may be mature polypeptides, as defined herein, and may or may not have an amino-terminal methionine residue, depending on the method by which they are prepared.

The term "IL-1ra-R polypeptide fragment" refers to a polypeptide that comprises a truncation at the amino-terminus (with or without a leader sequence) and/or a truncation at the carboxyl-terminus of the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. The term "IL-1ra-R polypeptide fragment" also refers to amino-terminal and/or carboxylterminal truncations of IL-1ra-R polypeptide orthologs, IL-1ra-R polypeptide derivatives, or IL-1ra-R polypeptide variants, or to amino-terminal and/or carboxyl-terminal truncations of the polypeptides encoded by IL-1ra-R polypeptide allelic variants or IL-1ra-R polypeptide splice variants. IL-1ra-R polypeptide fragments may result from alternative RNA splicing or from in vivo protease activity. Membrane-bound forms of an IL-1ra-R polypeptide are also contemplated by the present invention. In preferred embodiments, truncations and/or deletions comprise about 10 amino acids, or about 20 amino acids, or about 50 amino acids, or about 75 amino acids, or about 100 amino acids, or more than about 100 amino acids. The polypeptide fragments so produced will comprise about 25 contiguous amino acids, or about 50 amino acids, or about 75 amino acids, or about 100 amino acids, or about 125 amino acids. Such IL-1ra-R

10

15

20

polypeptide fragments may optionally comprise an amino-terminal methionine residue. It will be appreciated that such fragments can be used, for example, to generate antibodies to IL-1ra-R polypeptides.

The term "IL-1ra-R polypeptide ortholog" refers to a polypeptide from another species that corresponds to IL-1ra-R polypeptide amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. For example, mouse and human IL-1ra-R polypeptides are considered orthologs of each other.

The term "IL-1ra-R polypeptide variants" refers to IL-1ra-R polypeptides comprising amino acid sequences having one or more amino acid sequence substitutions, deletions (such as internal deletions and/or IL-1ra-R polypeptide fragments), and/or additions (such as internal additions and/or IL-1ra-R fusion polypeptides) as compared to the IL-1ra-R polypeptide amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 (with or without a leader sequence). Variants may be naturally occurring (e.g., IL-1ra-R polypeptide allelic variants, IL-1ra-R polypeptide orthologs, and IL-1ra-R polypeptide splice variants) or artificially constructed. Such IL-1ra-R polypeptide variants may be prepared from the corresponding nucleic acid molecules having a DNA sequence that varies accordingly from the DNA sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35. In preferred embodiments, the variants have from 1 to 3, or from 1 to 5, or from 1 to 10, or from 1 to 15, or from 1 to 20, or from 1 to 25, or from 1 to 50, or from 1 to 75, or from 1 to 100, or more than 100 amino acid substitutions, insertions, additions and/or deletions, wherein the substitutions may be conservative, or non-conservative, or any combination thereof.

The term "IL-1ra-R polypeptide derivatives" refers to the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, IL-1ra-R polypeptide fragments, IL-1ra-R polypeptide orthologs, or IL-1ra-R polypeptide variants, as defined herein, that have been chemically modified. The term "IL-1ra-R polypeptide derivatives" also refers to the polypeptides encoded

25

10

15

20

25

30

by IL-1ra-R polypeptide allelic variants or IL-1ra-R polypeptide splice variants, as defined herein, that have been chemically modified.

The term "mature IL-1ra-R polypeptide" refers to an IL-1ra-R polypeptide lacking a leader sequence. A mature IL-1ra-R polypeptide may also include other modifications such as proteolytic processing of the amino-terminus (with or without a leader sequence) and/or the carboxyl-terminus, cleavage of a smaller polypeptide from a larger precursor, N-linked and/or O-linked glycosylation, and the like.

The term "IL-1ra-R fusion polypeptide" refers to a fusion of one or more amino acids (such as a heterologous protein or peptide) at the amino- or carboxylterminus of the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, IL-1ra-R polypeptide fragments, IL-1ra-R polypeptide orthologs, IL-1ra-R polypeptide variants, or IL-1ra-R derivatives, as defined herein. The term "IL-1ra-R fusion polypeptide" also refers to a fusion of one or more amino acids at the amino- or carboxyl-terminus of the polypeptide encoded by IL-1ra-R polypeptide allelic variants or IL-1ra-R polypeptide splice variants, as defined herein.

The term "biologically active IL-1ra-R polypeptides" refers to IL-1ra-R polypeptides having at least one activity characteristic of the polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. In addition, an IL-1ra-R polypeptide may be active as an immunogen; that is, the IL-1ra-R polypeptide contains at least one epitope to which antibodies may be raised.

The term "isolated polypeptide" refers to a polypeptide of the present invention that (1) has been separated from at least about 50 percent of polynucleotides, lipids, carbohydrates, or other materials with which it is naturally found when isolated from the source cell, (2) is not linked (by covalent or noncovalent interaction) to all or a portion of a polypeptide to which the "isolated polypeptide" is linked in nature, (3) is operably linked (by covalent or noncovalent interaction) to a polypeptide with which it is not linked in nature, or (4) does not occur in nature. Preferably, the isolated polypeptide is substantially

10

15

20

25

30

free from any other contaminating polypeptides or other contaminants that are found in its natural environment that would interfere with its therapeutic, diagnostic, prophylactic or research use.

The term "identity," as known in the art, refers to a relationship between the sequences of two or more polypeptide molecules or two or more nucleic acid molecules, as determined by comparing the sequences. In the art, "identity" also means the degree of sequence relatedness between nucleic acid molecules or polypeptides, as the case may be, as determined by the match between strings of two or more nucleotide or two or more amino acid sequences. "Identity" measures the percent of identical matches between the smaller of two or more sequences with gap alignments (if any) addressed by a particular mathematical model or computer program (i.e., "algorithms").

The term "similarity" is a related concept, but in contrast to "identity," "similarity" refers to a measure of relatedness which includes both identical matches and conservative substitution matches. If two polypeptide sequences have, for example, 10/20 identical amino acids, and the remainder are all non-conservative substitutions, then the percent identity and similarity would both be 50%. If in the same example, there are five more positions where there are conservative substitutions, then the percent identity remains 50%, but the percent similarity would be 75% (15/20). Therefore, in cases where there are conservative substitutions, the percent similarity between two polypeptides will be higher than the percent identity between those two polypeptides.

The term "naturally occurring" or "native" when used in connection with biological materials such as nucleic acid molecules, polypeptides, host cells, and the like, refers to materials which are found in nature and are not manipulated by man. Similarly, "non-naturally occurring" or "non-native" as used herein refers to a material that is not found in nature or that has been structurally modified or synthesized by man.

The terms "effective amount" and "therapeutically effective amount" each refer to the amount of an IL-1ra-R polypeptide or IL-1ra-R nucleic acid molecule

10

15

20

25

30

used to support an observable level of one or more biological activities of the IL-1ra-R polypeptides as set forth herein.

The term "pharmaceutically acceptable carrier" or "physiologically acceptable carrier" as used herein refers to one or more formulation materials suitable for accomplishing or enhancing the delivery of the IL-1ra-R polypeptide, IL-1ra-R nucleic acid molecule, or IL-1ra-R selective binding agent as a pharmaceutical composition.

The term "antigen" refers to a molecule or a portion of a molecule capable of being bound by a selective binding agent, such as an antibody, and additionally capable of being used in an animal to produce antibodies capable of binding to an epitope of that antigen. An antigen may have one or more epitopes.

The term "selective binding agent" refers to a molecule or molecules having specificity for an IL-1ra-R polypeptide. As used herein, the terms, "specific" and "specificity" refer to the ability of the selective binding agents to bind to human IL-1ra-R polypeptides and not to bind to human non-IL-1ra-R polypeptides. It will be appreciated, however, that the selective binding agents may also bind orthologs of the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, that is, interspecies versions thereof, such as mouse and rat IL-1ra-R polypeptides.

The term "transduction" is used to refer to the transfer of genes from one bacterium to another, usually by a phage. "Transduction" also refers to the acquisition and transfer of eukaryotic cellular sequences by retroviruses.

The term "transfection" is used to refer to the uptake of foreign or exogenous DNA by a cell, and a cell has been "transfected" when the exogenous DNA has been introduced inside the cell membrane. A number of transfection techniques are well known in the art and are disclosed herein. See, e.g., Graham et al., 1973, Virology 52:456; Sambrook et al., Molecular Cloning, A Laboratory Manual (Cold Spring Harbor Laboratories, 1989); Davis et al., Basic Methods in Molecular Biology (Elsevier, 1986); and Chu et al., 1981, Gene 13:197. Such techniques can be used to introduce one or more exogenous DNA moieties into suitable host cells.

The term "transformation" as used herein refers to a change in a cell's genetic characteristics, and a cell has been transformed when it has been modified to contain a new DNA. For example, a cell is transformed where it is genetically modified from its native state. Following transfection or transduction, the transforming DNA may recombine with that of the cell by physically integrating into a chromosome of the cell, may be maintained transiently as an episomal element without being replicated, or may replicate independently as a plasmid. A cell is considered to have been stably transformed when the DNA is replicated with the division of the cell.

10

15

20

25

30

5

Relatedness of Nucleic Acid Molecules and/or Polypeptides

It is understood that related nucleic acid molecules include allelic or splice variants of the nucleic acid molecule of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, and include sequences which are complementary to any of the above nucleotide sequences. Related nucleic acid molecules also include a nucleotide sequence encoding a polypeptide comprising or consisting essentially of a substitution, modification, addition and/or deletion of one or more amino acid residues compared to the polypeptide in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. Such related IL-1ra-R polypeptides may comprise, for example, an addition and/or a deletion of one or more N-linked or O-linked glycosylation sites or an addition and/or a deletion of one or more cysteine residues.

Related nucleic acid molecules also include fragments of IL-1ra-R nucleic acid molecules which encode a polypeptide of at least about 25 contiguous amino acids, or about 50 amino acids, or about 75 amino acids, or about 100 amino acids, or about 125 amino acids, or more than 125 amino acid residues of the IL-1ra-R polypeptide of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

In addition, related IL-1ra-R nucleic acid molecules also include those molecules which comprise nucleotide sequences which hybridize under moderately or highly stringent conditions as defined herein with the fully

10

15

20

25

30

complementary sequence of the IL-1ra-R nucleic acid molecule of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, or of a molecule encoding a polypeptide, which polypeptide comprises the amino acid sequence as shown in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or of a nucleic acid fragment as defined herein, or of a nucleic acid fragment encoding a polypeptide as defined herein. Hybridization probes may be prepared using the IL-1ra-R sequences provided herein to screen cDNA, genomic or synthetic DNA libraries for related sequences. Regions of the DNA and/or amino acid sequence of IL-1ra-R polypeptide that exhibit significant identity to known sequences are readily determined using sequence alignment algorithms as described herein and those regions may be used to design probes for screening.

The term "highly stringent conditions" refers to those conditions that are designed to permit hybridization of DNA strands whose sequences are highly complementary, and to exclude hybridization of significantly mismatched DNAs. Hybridization stringency is principally determined by temperature, ionic strength, and the concentration of denaturing agents such as formamide. Examples of "highly stringent conditions" for hybridization and washing are 0.015 M sodium chloride, 0.0015 M sodium citrate at 65-68°C or 0.015 M sodium chloride, 0.0015 M sodium citrate, and 50% formamide at 42°C. See Sambrook, Fritsch & Maniatis, Molecular Cloning: A Laboratory Manual (2nd ed., Cold Spring Harbor Laboratory, 1989); Anderson et al., Nucleic Acid Hybridisation: A Practical Approach Ch. 4 (IRL Press Limited).

More stringent conditions (such as higher temperature, lower ionic strength, higher formamide, or other denaturing agent) may also be used – however, the rate of hybridization will be affected. Other agents may be included in the hybridization and washing buffers for the purpose of reducing non-specific and/or background hybridization. Examples are 0.1% bovine serum albumin, 0.1% polyvinyl-pyrrolidone, 0.1% sodium pyrophosphate, 0.1% sodium dodecylsulfate, NaDodSO₄, (SDS), ficoll, Denhardt's solution, sonicated salmon sperm DNA (or another non-complementary DNA), and dextran sulfate, although other suitable agents can also be used. The concentration and types of these

10

15

20

25

30

additives can be changed without substantially affecting the stringency of the hybridization conditions. Hybridization experiments are usually carried out at pH 6.8-7.4; however, at typical ionic strength conditions, the rate of hybridization is nearly independent of pH. See Anderson et al., Nucleic Acid Hybridisation: A Practical Approach Ch. 4 (IRL Press Limited).

Factors affecting the stability of DNA duplex include base composition, length, and degree of base pair mismatch. Hybridization conditions can be adjusted by one skilled in the art in order to accommodate these variables and allow DNAs of different sequence relatedness to form hybrids. The melting temperature of a perfectly matched DNA duplex can be estimated by the following equation:

 $T_m(^{\circ}C) = 81.5 + 16.6(log[Na+]) + 0.41(\%G+C) - 600/N - 0.72(\%formamide)$ where N is the length of the duplex formed, [Na+] is the molar concentration of the sodium ion in the hybridization or washing solution, %G+C is the percentage of (guanine+cytosine) bases in the hybrid. For imperfectly matched hybrids, the melting temperature is reduced by approximately $1^{\circ}C$ for each 1% mismatch.

The term "moderately stringent conditions" refers to conditions under which a DNA duplex with a greater degree of base pair mismatching than could occur under "highly stringent conditions" is able to form. Examples of typical "moderately stringent conditions" are 0.015 M sodium chloride, 0.0015 M sodium citrate at 50-65°C or 0.015 M sodium chloride, 0.0015 M sodium citrate, and 20% formamide at 37-50°C. By way of example, "moderately stringent conditions" of 50°C in 0.015 M sodium ion will allow about a 21% mismatch.

It will be appreciated by those skilled in the art that there is no absolute distinction between "highly stringent conditions" and "moderately stringent conditions." For example, at 0.015 M sodium ion (no formamide), the melting temperature of perfectly matched long DNA is about 71°C. With a wash at 65°C (at the same ionic strength), this would allow for approximately a 6% mismatch. To capture more distantly related sequences, one skilled in the art can simply lower the temperature or raise the ionic strength.

10

15

20

25

30

A good estimate of the melting temperature in 1M NaCl* for oligonucleotide probes up to about 20nt is given by:

 $Tm = 2^{\circ}C$ per A-T base pair + $4^{\circ}C$ per G-C base pair

*The sodium ion concentration in 6X salt sodium citrate (SSC) is 1M. See Suggs et al., Developmental Biology Using Purified Genes 683 (Brown and Fox, eds., 1981).

High stringency washing conditions for oligonucleotides are usually at a temperature of 0-5°C below the Tm of the oligonucleotide in 6X SSC, 0.1% SDS.

In another embodiment, related nucleic acid molecules comprise or consist of a nucleotide sequence that is at least about 70 percent identical to the nucleotide sequence as shown in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, or comprise or consist essentially of a nucleotide sequence encoding a polypeptide that is at least about 70 percent identical to the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. In preferred embodiments, the nucleotide sequences are about 75 percent, or about 80 percent, or about 85 percent, or about 90 percent, or about 95, 96, 97, 98, or 99 percent identical to the nucleotide sequence as shown in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, or the nucleotide sequences encode a polypeptide that is about 75 percent, or about 80 percent, or about 85 percent, or about 90 percent, or about 95, 96, 97, 98, or 99 percent identical to the polypeptide sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. Related nucleic acid molecules encode polypeptides possessing at least one activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

Differences in the nucleic acid sequence may result in conservative and/or non-conservative modifications of the amino acid sequence relative to the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

Conservative modifications to the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 (and the corresponding

10

15

20

modifications to the encoding nucleotides) will produce a polypeptide having functional and chemical characteristics similar to those of IL-1ra-R polypeptides. In contrast, substantial modifications in the functional and/or chemical characteristics of IL-1ra-R polypeptides may be accomplished by selecting substitutions in the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 that differ significantly in their effect on maintaining (a) the structure of the molecular backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain.

For example, a "conservative amino acid substitution" may involve a substitution of a native amino acid residue with a nonnative residue such that there is little or no effect on the polarity or charge of the amino acid residue at that position. Furthermore, any native residue in the polypeptide may also be substituted with alanine, as has been previously described for "alanine scanning mutagenesis."

Conservative amino acid substitutions also encompass non-naturally occurring amino acid residues that are typically incorporated by chemical peptide synthesis rather than by synthesis in biological systems. These include peptidomimetics, and other reversed or inverted forms of amino acid moieties.

Naturally occurring residues may be divided into classes based on common side chain properties:

- 1) hydrophobic: norleucine, Met, Ala, Val, Leu, Ile;
- 2) neutral hydrophilic: Cys, Ser, Thr;
- 3) acidic: Asp, Glu;
- 4) basic: Asn, Gln, His, Lys, Arg;
 - 5) residues that influence chain orientation: Gly, Pro; and
 - 6) aromatic: Trp, Tyr, Phe.

For example, non-conservative substitutions may involve the exchange of a member of one of these classes for a member from another class. Such substituted residues may be introduced into regions of the human IL-1ra-R

30

25

10

15

20

25

30

polypeptide that are homologous with non-human IL-1ra-R polypeptides, or into the non-homologous regions of the molecule.

In making such changes, the hydropathic index of amino acids may be considered. Each amino acid has been assigned a hydropathic index on the basis of its hydrophobicity and charge characteristics. The hydropathic indices are: isoleucine (+4.5); valine (+4.2); leucine (+3.8); phenylalanine (+2.8); cysteine/cystine (+2.5); methionine (+1.9); alanine (+1.8); glycine (-0.4); threonine (-0.7); serine (-0.8); tryptophan (-0.9); tyrosine (-1.3); proline (-1.6); histidine (-3.2); glutamate (-3.5); glutamine (-3.5); aspartate (-3.5); asparagine (-3.5); lysine (-3.9); and arginine (-4.5).

The importance of the hydropathic amino acid index in conferring interactive biological function on a protein is generally understood in the art (Kyte et al., 1982, J. Mol. Biol. 157:105-31). It is known that certain amino acids may be substituted for other amino acids having a similar hydropathic index or score and still retain a similar biological activity. In making changes based upon the hydropathic index, the substitution of amino acids whose hydropathic indices are within ± 2 is preferred, those which are within ± 1 are particularly preferred, and those within ± 0.5 are even more particularly preferred.

It is also understood in the art that the substitution of like amino acids can be made effectively on the basis of hydrophilicity, particularly where the biologically functionally equivalent protein or peptide thereby created is intended for use in immunological embodiments, as in the present case. The greatest local average hydrophilicity of a protein, as governed by the hydrophilicity of its adjacent amino acids, correlates with its immunogenicity and antigenicity, *i.e.*, with a biological property of the protein.

The following hydrophilicity values have been assigned to these amino acid residues: arginine (+3.0); lysine (+3.0); aspartate (+3.0 \pm 1); glutamate (+3.0 \pm 1); serine (+0.3); asparagine (+0.2); glutamine (+0.2); glycine (0); threonine (-0.4); proline (-0.5 \pm 1); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine (-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); and tryptophan (-3.4). In making changes based upon

similar hydrophilicity values, the substitution of amino acids whose hydrophilicity values are within ± 2 is preferred, those which are within ± 1 are particularly preferred, and those within ± 0.5 are even more particularly preferred. One may also identify epitopes from primary amino acid sequences on the basis of hydrophilicity. These regions are also referred to as "epitopic core regions."

Desired amino acid substitutions (whether conservative or non-conservative) can be determined by those skilled in the art at the time such substitutions are desired. For example, amino acid substitutions can be used to identify important residues of the IL-1ra-R polypeptide, or to increase or decrease the affinity of the IL-1ra-R polypeptides described herein. Exemplary amino acid substitutions are set forth in Table I.

<u>Table I</u>

Amino Acid Substitutions

Original Residues	Exemplary Substitutions	Preferred Substitutions
Ala	Val, Leu, Ile	Val
Arg	Lys, Gln, Asn	Lys
Asn	Gln	Gln
Asp	Glu	Glu
Cys	Ser, Ala	Ser
Gln	Asn	Asn
Glu	Asp	Asp
Gly	Pro, Ala	Ala
His	Asn, Gln, Lys, Arg	Arg
Ile	Leu, Val, Met, Ala,	Leu
	Phe, Norleucine	
Leu	Norleucine, Ile, Ile	
	Val, Met, Ala, Phe	
Lys	Arg, 1,4 Diamino-butyric	Arg
	Acid, Gln, Asn	

10

5

Met	Leu, Phe, Ile	Leu
Phe	Leu, Val, Ile, Ala,	Leu
	Tyr	
Pro	Ala	Gly
Ser	Thr, Ala, Cys	Thr
Thr	Ser	Ser
Trp	Tyr, Phe	Tyr
Tyr	Trp, Phe, Thr, Ser	Phe
Val	Ile, Met, Leu, Phe,	Leu
	Ala, Norleucine	

A skilled artisan will be able to determine suitable variants of the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 using well-known techniques. For identifying suitable areas of the molecule that may be changed without destroying biological activity, one skilled in the art may target areas not believed to be important for activity. For example, when similar polypeptides with similar activities from the same species or from other species are known, one skilled in the art may compare the amino acid sequence of an IL-1ra-R polypeptide to such similar polypeptides. With such a comparison, one can identify residues and portions of the molecules that are conserved among similar polypeptides. It will be appreciated that changes in areas of the IL-1ra-R molecule that are not conserved relative to such similar polypeptides would be less likely to adversely affect the biological activity and/or structure of an IL-1ra-R polypeptide. One skilled in the art would also know that, even in relatively conserved regions, one may substitute chemically similar amino acids for the naturally occurring residues while retaining activity (conservative amino acid residue substitutions). Therefore, even areas that may be important for biological activity or for structure may be subject to conservative amino acid substitutions without destroying the biological activity or without adversely affecting the polypeptide structure.

20

10

15

10

15

20

25

30

Additionally, one skilled in the art can review structure-function studies identifying residues in similar polypeptides that are important for activity or structure. In view of such a comparison, one can predict the importance of amino acid residues in an IL-1ra-R polypeptide that correspond to amino acid residues that are important for activity or structure in similar polypeptides. One skilled in the art may opt for chemically similar amino acid substitutions for such predicted important amino acid residues of IL-1ra-R polypeptides.

One skilled in the art can also analyze the three-dimensional structure and amino acid sequence in relation to that structure in similar polypeptides. In view of such information, one skilled in the art may predict the alignment of amino acid residues of IL-1ra-R polypeptide with respect to its three dimensional structure. One skilled in the art may choose not to make radical changes to amino acid residues predicted to be on the surface of the protein, since such residues may be involved in important interactions with other molecules. Moreover, one skilled in the art may generate test variants containing a single amino acid substitution at each amino acid residue. The variants could be screened using activity assays known to those with skill in the art. Such variants could be used to gather information about suitable variants. For example, if one discovered that a change to a particular amino acid residue resulted in destroyed, undesirably reduced, or unsuitable activity, variants with such a change would be avoided. In other words, based on information gathered from such routine experiments, one skilled in the art can readily determine the amino acids where further substitutions should be avoided either alone or in combination with other mutations.

A number of scientific publications have been devoted to the prediction of secondary structure. See Moult, 1996, Curr. Opin. Biotechnol. 7:422-27; Chou et al., 1974, Biochemistry 13:222-45; Chou et al., 1974, Biochemistry 113:211-22; Chou et al., 1978, Adv. Enzymol. Relat. Areas Mol. Biol. 47:45-48; Chou et al., 1978, Ann. Rev. Biochem. 47:251-276; and Chou et al., 1979, Biophys. J. 26:367-84. Moreover, computer programs are currently available to assist with predicting secondary structure. One method of predicting secondary structure is based upon homology modeling. For example, two polypeptides or proteins which have a

10

15

20

25

30

sequence identity of greater than 30%, or similarity greater than 40%, often have similar structural topologies. The recent growth of the protein structural database (PDB) has provided enhanced predictability of secondary structure, including the potential number of folds within the structure of a polypeptide or protein. *See* Holm *et al.*, 1999, *Nucleic Acids Res.* 27:244-47. It has been suggested that there are a limited number of folds in a given polypeptide or protein and that once a critical number of structures have been resolved, structural prediction will become dramatically more accurate (Brenner *et al.*, 1997, *Curr. Opin. Struct. Biol.* 7:369-76).

Additional methods of predicting secondary structure include "threading" (Jones, 1997, Curr. Opin. Struct. Biol. 7:377-87; Sippl et al., 1996, Structure 4:15-19), "profile analysis" (Bowie et al., 1991, Science, 253:164-70; Gribskov et al., 1990, Methods Enzymol. 183:146-59; Gribskov et al., 1987, Proc. Nat. Acad. Sci. U.S.A. 84:4355-58), and "evolutionary linkage" (See Holm et al., supra, and Brenner et al., supra).

Preferred IL-1ra-R polypeptide variants include glycosylation variants wherein the number and/or type of glycosylation sites have been altered compared to the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. In one embodiment, IL-1ra-R polypeptide variants comprise a greater or a lesser number of N-linked glycosylation sites than the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. An N-linked glycosylation site is characterized by the sequence: Asn-X-Ser or Asn-X-Thr, wherein the amino acid residue designated as X may be any amino acid residue except proline. The substitution of amino acid residues to create this sequence provides a potential new site for the addition of an N-linked carbohydrate chain. Alternatively, substitutions that eliminate this sequence will remove an existing N-linked carbohydrate chain. Also provided is a rearrangement of N-linked carbohydrate chains wherein one or more N-linked glycosylation sites (typically those that are naturally occurring) are eliminated and one or more new N-linked sites are created. Additional preferred IL-1ra-R variants include cysteine variants, wherein one or more cysteine residues

10

15

20

are deleted or substituted with another amino acid (e.g., serine) as compared to the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. Cysteine variants are useful when IL-1ra-R polypeptides must be refolded into a biologically active conformation such as after the isolation of insoluble inclusion bodies. Cysteine variants generally have fewer cysteine residues than the native protein, and typically have an even number to minimize interactions resulting from unpaired cysteines.

In other embodiments, related nucleic acid molecules comprise or consist of a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid insertion and wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid deletion and wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. Related nucleic acid molecules also comprise or consist of a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 wherein the polypeptide has a carboxyl- and/or amino-terminal truncation and further wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36. Related nucleic acid molecules also comprise or consist of a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one modification selected from the group consisting of amino acid substitutions, amino acid insertions, amino acid deletions, carboxyl-terminal truncations, and aminoterminal truncations and wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

In addition, the polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-

30

25

10

15

20

25

30

Ira-R polypeptide, may be fused to a homologous polypeptide to form a homodimer or to a heterologous polypeptide to form a heterodimer. Heterologous peptides and polypeptides include, but are not limited to: an epitope to allow for the detection and/or isolation of an IL-1ra-R fusion polypeptide; a transmembrane receptor protein or a portion thereof, such as an extracellular domain or a transmembrane and intracellular domain; a ligand or a portion thereof which binds to a transmembrane receptor protein; an enzyme or portion thereof which is catalytically active; a polypeptide or peptide which promotes oligomerization, such as a leucine zipper domain; a polypeptide or peptide which increases stability, such as an immunoglobulin constant region; and a polypeptide which has a therapeutic activity different from the polypeptide comprising the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-1ra-R polypeptide.

Fusions can be made either at the amino-terminus or at the carboxyl-terminus of the polypeptide comprising the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-1ra-R polypeptide. Fusions may be direct with no linker or adapter molecule or may be through a linker or adapter molecule. A linker or adapter molecule may be one or more amino acid residues, typically from about 20 to about 50 amino acid residues. A linker or adapter molecule may also be designed with a cleavage site for a DNA restriction endonuclease or for a protease to allow for the separation of the fused moieties. It will be appreciated that once constructed, the fusion polypeptides can be derivatized according to the methods described herein.

In a further embodiment of the invention, the polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-1ra-R polypeptide, is fused to one or more domains of an Fc region of human IgG. Antibodies comprise two functionally independent parts, a variable domain known as "Fab," that binds an antigen, and a constant domain known as "Fc," that is involved in effector functions such as complement activation and attack by phagocytic cells. An Fc has a long serum half-life, whereas an Fab is short-lived. Capon *et al.*, 1989, *Nature* 337:525-31. When

constructed together with a therapeutic protein, an Fc domain can provide longer half-life or incorporate such functions as Fc receptor binding, protein A binding, complement fixation, and perhaps even placental transfer. *Id.* Table II summarizes the use of certain Fc fusions known in the art.

5

<u>Table II</u>
Fc Fusion with Therapeutic Proteins

Form of Fc	Fusion partner	Therapeutic implications	Reference
IgG1	N-terminus of CD30-L	Hodgkin's disease; anaplastic lymphoma; T- cell leukemia	U.S. Patent No. 5,480,981
Murine Fc&a	IL-10	anti-inflammatory; transplant rejection	Zheng et al., 1995, J. Immunol. 154:5590-600
IgG1	TNF receptor	septic shock	Fisher et al., 1996, N. Engl. J. Med. 334:1697- 1702; Van Zee et al., 1996, J. Immunol. 156:2221-30
IgG, IgA, IgM, or IgE (excluding the first domain)	TNF receptor	inflammation, autoimmune disorders	U.S. Patent No. 5,808,029
IgG1	CD4 receptor	AIDS	Capon et al., 1989, Nature 337: 525-31
IgG1, IgG3	N-terminus of IL-2	anti-cancer, antiviral	Harvill et al., 1995, Immunotech. 1:95-105
IgG1	C-terminus of OPG	osteoarthritis; bone density	WO 97/23614
IgG1	N-terminus of leptin	anti-obesity	PCT/US 97/23183, filed December 11, 1997
Human Ig C91	CTLA-4	autoimmune disorders	Linsley, 1991, <i>J. Exp. Med.</i> , 174:561-69

In one example, a human IgG hinge, CH2, and CH3 region may be fused at either the amino-terminus or carboxyl-terminus of the IL-1ra-R polypeptides using methods known to the skilled artisan. In another example, a human IgG hinge, CH2, and CH3 region may be fused at either the amino-terminus or carboxyl-terminus of an IL-1ra-R polypeptide fragment (e.g., the predicted extracellular portion of IL-1ra-R polypeptide).

The resulting IL-1ra-R fusion polypeptide may be purified by use of a Protein A affinity column. Peptides and proteins fused to an Fc region have been found to exhibit a substantially greater half-life *in vivo* than the unfused counterpart. Also, a fusion to an Fc region allows for dimerization/multimerization of the fusion polypeptide. The Fc region may be a naturally occurring Fc region, or may be altered to improve certain qualities, such as therapeutic qualities, circulation time, or reduced aggregation.

Identity and similarity of related nucleic acid molecules and polypeptides are readily calculated by known methods. Such methods include, but are not limited to those described in *Computational Molecular Biology* (A.M. Lesk, ed., Oxford University Press 1988); *Biocomputing: Informatics and Genome Projects* (D.W. Smith, ed., Academic Press 1993); *Computer Analysis of Sequence Data* (Part 1, A.M. Griffin and H.G. Griffin, eds., Humana Press 1994); G. von Heinle, *Sequence Analysis in Molecular Biology* (Academic Press 1987); *Sequence Analysis Primer* (M. Gribskov and J. Devereux, eds., M. Stockton Press 1991); and Carillo *et al.*, 1988, *SIAM J. Applied Math.*, 48:1073.

Preferred methods to determine identity and/or similarity are designed to give the largest match between the sequences tested. Methods to determine identity and similarity are described in publicly available computer programs. Preferred computer program methods to determine identity and similarity between two sequences include, but are not limited to, the GCG program package, including GAP (Devereux et al., 1984, Nucleic Acids Res. 12:387; Genetics Computer Group, University of Wisconsin, Madison, WI), BLASTP, BLASTN, and FASTA (Altschul et al., 1990, J. Mol. Biol. 215:403-10). The BLASTX program is publicly available from the National Center for Biotechnology Information (NCBI) and other sources (Altschul et al., BLAST Manual (NCB NLM NIH, Bethesda, MD); Altschul et al., 1990, supra). The well-known Smith Waterman algorithm may also be used to determine identity.

Certain alignment schemes for aligning two amino acid sequences may result in the matching of only a short region of the two sequences, and this small aligned region may have very high sequence identity even though there is no

25

30

5

10

15

10

15

20

30

significant relationship between the two full-length sequences. Accordingly, in a preferred embodiment, the selected alignment method (GAP program) will result in an alignment that spans at least 50 contiguous amino acids of the claimed polypeptide.

For example, using the computer algorithm GAP (Genetics Computer Group, University of Wisconsin, Madison, WI), two polypeptides for which the percent sequence identity is to be determined are aligned for optimal matching of their respective amino acids (the "matched span," as determined by the algorithm). A gap opening penalty (which is calculated as 3X the average diagonal; the "average diagonal" is the average of the diagonal of the comparison matrix being used; the "diagonal" is the score or number assigned to each perfect amino acid match by the particular comparison matrix) and a gap extension penalty (which is usually 0.1X the gap opening penalty), as well as a comparison matrix such as PAM 250 or BLOSUM 62 are used in conjunction with the algorithm. A standard comparison matrix is also used by the algorithm (see Dayhoff et al., 5 Atlas of Protein Sequence and Structure (Supp. 3 1978)(PAM250 comparison matrix); Henikoff et al., 1992, Proc. Natl. Acad. Sci USA 89:10915-19 (BLOSUM 62 comparison matrix)).

Preferred parameters for polypeptide sequence comparison include the following:

Algorithm: Needleman and Wunsch, 1970, J. Mol. Biol. 48:443-53;

Comparison matrix: BLOSUM 62 (Henikoff et al., supra);

Gap Penalty: 12

Gap Length Penalty: 4

Threshold of Similarity: 0

The GAP program is useful with the above parameters. The aforementioned parameters are the default parameters for polypeptide comparisons (along with no penalty for end gaps) using the GAP algorithm.

Preferred parameters for nucleic acid molecule sequence comparison include the following:

Algorithm: Needleman and Wunsch, supra;

Comparison matrix: matches = +10, mismatch = 0

Gap Penalty: 50

Gap Length Penalty: 3

The GAP program is also useful with the above parameters. The aforementioned parameters are the default parameters for nucleic acid molecule comparisons.

Other exemplary algorithms, gap opening penalties, gap extension penalties, comparison matrices, and thresholds of similarity may be used, including those set forth in the Program Manual, Wisconsin Package, Version 9, September, 1997. The particular choices to be made will be apparent to those of skill in the art and will depend on the specific comparison to be made, such as DNA-to-DNA, protein-to-protein, protein-to-DNA; and additionally, whether the comparison is between given pairs of sequences (in which case GAP or BestFit are generally preferred) or between one sequence and a large database of sequences (in which case FASTA or BLASTA are preferred).

20

25

30

5

10

15

Nucleic Acid Molecules

The nucleic acid molecules encoding a polypeptide comprising the amino acid sequence of an IL-1ra-R polypeptide can readily be obtained in a variety of ways including, without limitation, chemical synthesis, cDNA or genomic library screening, expression library screening, and/or PCR amplification of cDNA.

Recombinant DNA methods used herein are generally those set forth in Sambrook et al., Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 1989) and/or Current Protocols in Molecular Biology (Ausubel et al., eds., Green Publishers Inc. and Wiley and Sons 1994). The invention provides for nucleic acid molecules as described herein and methods for obtaining such molecules.

10

15

20

25

30

Where a gene encoding the amino acid sequence of an IL-1ra-R polypeptide has been identified from one species, all or a portion of that gene may be used as a probe to identify orthologs or related genes from the same species. The probes or primers may be used to screen cDNA libraries from various tissue sources believed to express the IL-1ra-R polypeptide. In addition, part or all of a nucleic acid molecule having the sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35 may be used to screen a genomic library to identify and isolate a gene encoding the amino acid sequence of an IL-1ra-R polypeptide. Typically, conditions of moderate or high stringency will be employed for screening to minimize the number of false positives obtained from the screening.

Nucleic acid molecules encoding the amino acid sequence of IL-1ra-R polypeptides may also be identified by expression cloning which employs the detection of positive clones based upon a property of the expressed protein. Typically, nucleic acid libraries are screened by the binding an antibody or other binding partner (e.g., receptor or ligand) to cloned proteins that are expressed and displayed on a host cell surface. The antibody or binding partner is modified with a detectable label to identify those cells expressing the desired clone.

Recombinant expression techniques conducted in accordance with the descriptions set forth below may be followed to produce these polynucleotides and to express the encoded polypeptides. For example, by inserting a nucleic acid sequence that encodes the amino acid sequence of an IL-1ra-R polypeptide into an appropriate vector, one skilled in the art can readily produce large quantities of the desired nucleotide sequence. The sequences can then be used to generate detection probes or amplification primers. Alternatively, a polynucleotide encoding the amino acid sequence of an IL-1ra-R polypeptide can be inserted into an expression vector. By introducing the expression vector into an appropriate host, the encoded IL-1ra-R polypeptide may be produced in large amounts.

Another method for obtaining a suitable nucleic acid sequence is the polymerase chain reaction (PCR). In this method, cDNA is prepared from poly(A)+RNA or total RNA using the enzyme reverse transcriptase. Two

10

15

20

25

30

primers, typically complementary to two separate regions of cDNA encoding the amino acid sequence of an IL-1ra-R polypeptide, are then added to the cDNA along with a polymerase such as *Taq* polymerase, and the polymerase amplifies the cDNA region between the two primers.

Another means of preparing a nucleic acid molecule encoding the amino acid sequence of an IL-1ra-R polypeptide is chemical synthesis using methods well known to the skilled artisan such as those described by Engels et al., 1989, Angew. Chem. Intl. Ed. 28:716-34. These methods include, inter alia, the phosphotriester, phosphoramidite, and H-phosphonate methods for nucleic acid synthesis. A preferred method for such chemical synthesis is polymer-supported synthesis using standard phosphoramidite chemistry. Typically, the DNA encoding the amino acid sequence of an IL-1ra-R polypeptide will be several hundred nucleotides in length. Nucleic acids larger than about 100 nucleotides can be synthesized as several fragments using these methods. The fragments can then be ligated together to form the full-length nucleotide sequence of an IL-1ra-R Usually, the DNA fragment encoding the amino-terminus of the gene. polypeptide will have an ATG, which encodes a methionine residue. methionine may or may not be present on the mature form of the IL-1ra-R polypeptide, depending on whether the polypeptide produced in the host cell is designed to be secreted from that cell. Other methods known to the skilled artisan may be used as well.

In certain embodiments, nucleic acid variants contain codons which have been altered for optimal expression of an IL-1ra-R polypeptide in a given host cell. Particular codon alterations will depend upon the IL-1ra-R polypeptide and host cell selected for expression. Such "codon optimization" can be carried out by a variety of methods, for example, by selecting codons which are preferred for use in highly expressed genes in a given host cell. Computer algorithms which incorporate codon frequency tables such as "Eco_high.Cod" for codon preference of highly expressed bacterial genes may be used and are provided by the University of Wisconsin Package Version 9.0 (Genetics Computer Group, frequency tables include useful codon WI). Other Madison,

10

15

20

"Celegans_high.cod," "Celegans_low.cod," "Drosophila_high.cod," "Human_high.cod," "Maize_high.cod," and "Yeast_high.cod."

In some cases, it may be desirable to prepare nucleic acid molecules encoding IL-1ra-R polypeptide variants. Nucleic acid molecules encoding variants may be produced using site directed mutagenesis, PCR amplification, or other appropriate methods, where the primer(s) have the desired point mutations (see Sambrook et al., supra, and Ausubel et al., supra, for descriptions of mutagenesis techniques). Chemical synthesis using methods described by Engels et al., supra, may also be used to prepare such variants. Other methods known to the skilled artisan may be used as well.

Vectors and Host Cells

A nucleic acid molecule encoding the amino acid sequence of an IL-1ra-R polypeptide is inserted into an appropriate expression vector using standard ligation techniques. The vector is typically selected to be functional in the particular host cell employed (*i.e.*, the vector is compatible with the host cell machinery such that amplification of the gene and/or expression of the gene can occur). A nucleic acid molecule encoding the amino acid sequence of an IL-1ra-R polypeptide may be amplified/expressed in prokaryotic, yeast, insect (baculovirus systems) and/or eukaryotic host cells. Selection of the host cell will depend in part on whether an IL-1ra-R polypeptide is to be post-translationally modified (*e.g.*, glycosylated and/or phosphorylated). If so, yeast, insect, or mammalian host cells are preferable. For a review of expression vectors, *see Meth. Enz.*, vol. 185 (D.V. Goeddel, ed., Academic Press 1990).

Typically, expression vectors used in any of the host cells will contain sequences for plasmid maintenance and for cloning and expression of exogenous nucleotide sequences. Such sequences, collectively referred to as "flanking sequences" in certain embodiments will typically include one or more of the following nucleotide sequences: a promoter, one or more enhancer sequences, an origin of replication, a transcriptional termination sequence, a complete intron sequence containing a donor and acceptor splice site, a sequence encoding a

30

leader sequence for polypeptide secretion, a ribosome binding site, a polyadenylation sequence, a polylinker region for inserting the nucleic acid encoding the polypeptide to be expressed, and a selectable marker element. Each of these sequences is discussed below.

Optionally, the vector may contain a "tag"-encoding sequence, *i.e.*, an oligonucleotide molecule located at the 5' or 3' end of the IL-1ra-R polypeptide coding sequence; the oligonucleotide sequence encodes polyHis (such as hexaHis), or another "tag" such as FLAG, HA (hemaglutinin influenza virus), or *myc* for which commercially available antibodies exist. This tag is typically fused to the polypeptide upon expression of the polypeptide, and can serve as a means for affinity purification of the IL-1ra-R polypeptide from the host cell. Affinity purification can be accomplished, for example, by column chromatography using antibodies against the tag as an affinity matrix. Optionally, the tag can subsequently be removed from the purified IL-1ra-R polypeptide by various means such as using certain peptidases for cleavage.

Flanking sequences may be homologous (*i.e.*, from the same species and/or strain as the host cell), heterologous (*i.e.*, from a species other than the host cell species or strain), hybrid (*i.e.*, a combination of flanking sequences from more than one source), or synthetic, or the flanking sequences may be native sequences which normally function to regulate IL-1ra-R polypeptide expression. As such, the source of a flanking sequence may be any prokaryotic or eukaryotic organism, any vertebrate or invertebrate organism, or any plant, provided that the flanking sequence is functional in, and can be activated by, the host cell machinery.

Flanking sequences useful in the vectors of this invention may be obtained by any of several methods well known in the art. Typically, flanking sequences useful herein – other than the IL-1ra-R gene flanking sequences – will have been previously identified by mapping and/or by restriction endonuclease digestion and can thus be isolated from the proper tissue source using the appropriate restriction endonucleases. In some cases, the full nucleotide sequence of a flanking

25

30

5

10

15

10

15

20

25

30

sequence may be known. Here, the flanking sequence may be synthesized using the methods described herein for nucleic acid synthesis or cloning.

Where all or only a portion of the flanking sequence is known, it may be obtained using PCR and/or by screening a genomic library with a suitable oligonucleotide and/or flanking sequence fragment from the same or another species. Where the flanking sequence is not known, a fragment of DNA containing a flanking sequence may be isolated from a larger piece of DNA that may contain, for example, a coding sequence or even another gene or genes. Isolation may be accomplished by restriction endonuclease digestion to produce the proper DNA fragment followed by isolation using agarose gel purification, Qiagen® column chromatography (Chatsworth, CA), or other methods known to the skilled artisan. The selection of suitable enzymes to accomplish this purpose will be readily apparent to one of ordinary skill in the art.

An origin of replication is typically a part of those prokaryotic expression vectors purchased commercially, and the origin aids in the amplification of the vector in a host cell. Amplification of the vector to a certain copy number can, in some cases, be important for the optimal expression of an IL-1ra-R polypeptide. If the vector of choice does not contain an origin of replication site, one may be chemically synthesized based on a known sequence, and ligated into the vector. For example, the origin of replication from the plasmid pBR322 (New England Biolabs, Beverly, MA) is suitable for most gram-negative bacteria and various origins (e.g., SV40, polyoma, adenovirus, vesicular stomatitus virus (VSV), or papillomaviruses such as HPV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (for example, the SV40 origin is often used only because it contains the early promoter).

A transcription termination sequence is typically located 3' of the end of a polypeptide coding region and serves to terminate transcription. Usually, a transcription termination sequence in prokaryotic cells is a G-C rich fragment followed by a poly-T sequence. While the sequence is easily cloned from a library or even purchased commercially as part of a vector, it can also be readily

10

15

20

synthesized using methods for nucleic acid synthesis such as those described herein.

A selectable marker gene element encodes a protein necessary for the survival and growth of a host cell grown in a selective culture medium. Typical selection marker genes encode proteins that (a) confer resistance to antibiotics or other toxins, *e.g.*, ampicillin, tetracycline, or kanamycin for prokaryotic host cells; (b) complement auxotrophic deficiencies of the cell; or (c) supply critical nutrients not available from complex media. Preferred selectable markers are the kanamycin resistance gene, the ampicillin resistance gene, and the tetracycline resistance gene. A neomycin resistance gene may also be used for selection in prokaryotic and eukaryotic host cells.

Other selection genes may be used to amplify the gene that will be expressed. Amplification is the process wherein genes that are in greater demand for the production of a protein critical for growth are reiterated in tandem within the chromosomes of successive generations of recombinant cells. Examples of suitable selectable markers for mammalian cells include dihydrofolate reductase (DHFR) and thymidine kinase. The mammalian cell transformants are placed under selection pressure wherein only the transformants are uniquely adapted to survive by virtue of the selection gene present in the vector. Selection pressure is imposed by culturing the transformed cells under conditions in which the concentration of selection agent in the medium is successively changed, thereby leading to the amplification of both the selection gene and the DNA that encodes an IL-1ra-R polypeptide. As a result, increased quantities of IL-1ra-R polypeptide are synthesized from the amplified DNA.

A ribosome binding site is usually necessary for translation initiation of mRNA and is characterized by a Shine-Dalgarno sequence (prokaryotes) or a Kozak sequence (eukaryotes). The element is typically located 3' to the promoter and 5' to the coding sequence of an IL-1ra-R polypeptide to be expressed. The Shine-Dalgarno sequence is varied but is typically a polypurine (*i.e.*, having a high A-G content). Many Shine-Dalgarno sequences have been identified, each

10

15

20

25

of which can be readily synthesized using methods set forth herein and used in a prokaryotic vector.

A leader, or signal, sequence may be used to direct an IL-1ra-R polypeptide out of the host cell. Typically, a nucleotide sequence encoding the signal sequence is positioned in the coding region of an IL-1ra-R nucleic acid molecule, or directly at the 5' end of an IL-1ra-R polypeptide coding region. Many signal sequences have been identified, and any of those that are functional in the selected host cell may be used in conjunction with an IL-1ra-R nucleic acid molecule. Therefore, a signal sequence may be homologous (naturally occurring) or heterologous to the IL-1ra-R nucleic acid molecule. Additionally, a signal sequence may be chemically synthesized using methods described herein. In most cases, the secretion of an IL-1ra-R polypeptide from the host cell via the presence of a signal peptide will result in the removal of the signal peptide from the secreted IL-1ra-R polypeptide. The signal sequence may be a component of the vector, or it may be a part of an IL-1ra-R nucleic acid molecule that is inserted into the vector.

Included within the scope of this invention is the use of either a nucleotide sequence encoding a native IL-1ra-R polypeptide signal sequence joined to an IL-1ra-R polypeptide coding region or a nucleotide sequence encoding a heterologous signal sequence joined to an IL-1ra-R polypeptide coding region. The heterologous signal sequence selected should be one that is recognized and processed, *i.e.*, cleaved by a signal peptidase, by the host cell. For prokaryotic host cells that do not recognize and process the native IL-1ra-R polypeptide signal sequence, the signal sequence is substituted by a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, or heat-stable enterotoxin II leaders. For yeast secretion, the native IL-1ra-R polypeptide signal sequence may be substituted by the yeast invertase, alpha factor, or acid phosphatase leaders. In mammalian cell expression the native signal sequence is satisfactory, although other mammalian signal sequences may

30 be suitable.

10

15

20

In some cases, such as where glycosylation is desired in a eukaryotic host cell expression system, one may manipulate the various presequences to improve glycosylation or yield. For example, one may alter the peptidase cleavage site of a particular signal peptide, or add pro-sequences, which also may affect glycosylation. The final protein product may have, in the -1 position (relative to the first amino acid of the mature protein) one or more additional amino acids incident to expression, which may not have been totally removed. For example, the final protein product may have one or two amino acid residues found in the peptidase cleavage site, attached to the amino-terminus. Alternatively, use of some enzyme cleavage sites may result in a slightly truncated form of the desired IL-1ra-R polypeptide, if the enzyme cuts at such area within the mature polypeptide.

In many cases, transcription of a nucleic acid molecule is increased by the presence of one or more introns in the vector; this is particularly true where a polypeptide is produced in eukaryotic host cells, especially mammalian host cells. The introns used may be naturally occurring within the IL-1ra-R gene especially where the gene used is a full-length genomic sequence or a fragment thereof. Where the intron is not naturally occurring within the gene (as for most cDNAs), the intron may be obtained from another source. The position of the intron with respect to flanking sequences and the IL-1ra-R gene is generally important, as the intron must be transcribed to be effective. Thus, when an IL-1ra-R cDNA molecule is being transcribed, the preferred position for the intron is 3' to the transcription start site and 5' to the poly-A transcription termination sequence. Preferably, the intron or introns will be located on one side or the other (i.e., 5' or 3') of the cDNA such that it does not interrupt the coding sequence. Any intron from any source, including viral, prokaryotic and eukaryotic (plant or animal) organisms, may be used to practice this invention, provided that it is compatible with the host cell into which it is inserted. Also included herein are synthetic introns. Optionally, more than one intron may be used in the vector.

The expression and cloning vectors of the present invention will typically contain a promoter that is recognized by the host organism and operably linked to

30

10

15

20

25

30

the molecule encoding the IL-1ra-R polypeptide. Promoters are untranscribed sequences located upstream (i.e., 5') to the start codon of a structural gene (generally within about 100 to 1000 bp) that control the transcription of the structural gene. Promoters are conventionally grouped into one of two classes: inducible promoters and constitutive promoters. Inducible promoters initiate increased levels of transcription from DNA under their control in response to some change in culture conditions, such as the presence or absence of a nutrient or a change in temperature. Constitutive promoters, on the other hand, initiate continual gene product production; that is, there is little or no control over gene expression. A large number of promoters, recognized by a variety of potential host cells, are well known. A suitable promoter is operably linked to the DNA encoding IL-1ra-R polypeptide by removing the promoter from the source DNA by restriction enzyme digestion and inserting the desired promoter sequence into The native IL-1ra-R promoter sequence may be used to direct the vector. amplification and/or expression of an IL-1ra-R nucleic acid molecule. heterologous promoter is preferred, however, if it permits greater transcription and higher yields of the expressed protein as compared to the native promoter, and if it is compatible with the host cell system that has been selected for use.

Promoters suitable for use with prokaryotic hosts include the betalactamase and lactose promoter systems; alkaline phosphatase; a tryptophan (trp) promoter system; and hybrid promoters such as the tac promoter. Other known bacterial promoters are also suitable. Their sequences have been published, thereby enabling one skilled in the art to ligate them to the desired DNA sequence, using linkers or adapters as needed to supply any useful restriction sites.

Suitable promoters for use with yeast hosts are also well known in the art. Yeast enhancers are advantageously used with yeast promoters. Suitable promoters for use with mammalian host cells are well known and include, but are not limited to, those obtained from the genomes of viruses such as polyoma virus, fowlpox virus, adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, retroviruses, hepatitis-B virus and most preferably Simian Virus 40 (SV40). Other suitable mammalian promoters include

10

15

20

25

30

heterologous mammalian promoters, for example, heat-shock promoters and the actin promoter.

Additional promoters which may be of interest in controlling IL-1ra-R gene expression include, but are not limited to: the SV40 early promoter region (Bernoist and Chambon, 1981, Nature 290:304-10); the CMV promoter; the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, et al., 1980, Cell 22:787-97); the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1444-45); the regulatory sequences of the metallothionine gene (Brinster et al., 1982, Nature 296:39-42); prokaryotic expression vectors such as the beta-lactamase promoter (Villa-Kamaroff et al., 1978, Proc. Natl. Acad. Sci. U.S.A., 75:3727-31); or the tac promoter (DeBoer et al., 1983, Proc. Natl. Acad. Sci. U.S.A., 80:21-25). Also of interest are the following animal transcriptional control regions, which exhibit tissue specificity and have been utilized in transgenic animals: the elastase I gene control region which is active in pancreatic acinar cells (Swift et al., 1984, Cell 38:639-46; Ornitz et al., 1986, Cold Spring Harbor Symp. Quant. Biol. 50:399-409 (1986); MacDonald, 1987, Hepatology 7:425-515); the insulin gene control region which is active in pancreatic beta cells (Hanahan, 1985, Nature 315:115-22); the immunoglobulin gene control region which is active in lymphoid cells (Grosschedl et al., 1984, Cell 38:647-58; Adames et al., 1985, Nature 318:533-38; Alexander et al., 1987, Mol. Cell. Biol., 7:1436-44); the mouse mammary tumor virus control region which is active in testicular, breast, lymphoid and mast cells (Leder et al., 1986, Cell 45:485-95); the albumin gene control region which is active in liver (Pinkert et al., 1987, Genes and Devel. 1:268-76); the alpha-fetoprotein gene control region which is active in liver (Krumlauf et al., 1985, Mol. Cell. Biol., 5:1639-48; Hammer et al., 1987, Science 235:53-58); the alpha 1antitrypsin gene control region which is active in the liver (Kelsey et al., 1987, Genes and Devel. 1:161-71); the beta-globin gene control region which is active in myeloid cells (Mogram et al., 1985, Nature 315:338-40; Kollias et al., 1986, Cell 46:89-94); the myelin basic protein gene control region which is active in oligodendrocyte cells in the brain (Readhead et al., 1987, Cell 48:703-12); the

10

15

20

25

30

myosin light chain-2 gene control region which is active in skeletal muscle (Sani, 1985, *Nature* 314:283-86); and the gonadotropic releasing hormone gene control region which is active in the hypothalamus (Mason *et al.*, 1986, *Science* 234:1372-78).

An enhancer sequence may be inserted into the vector to increase the transcription of a DNA encoding an IL-1ra-R polypeptide of the present invention by higher eukaryotes. Enhancers are cis-acting elements of DNA, usually about 10-300 bp in length, that act on the promoter to increase transcription. Enhancers are relatively orientation and position independent. They have been found 5' and 3' to the transcription unit. Several enhancer sequences available from mammalian genes are known (e.g., globin, elastase, albumin, alpha-feto-protein and insulin). Typically, however, an enhancer from a virus will be used. The SV40 enhancer, the cytomegalovirus early promoter enhancer, the polyoma enhancer, and adenovirus enhancers are exemplary enhancing elements for the activation of eukaryotic promoters. While an enhancer may be spliced into the vector at a position 5' or 3' to an IL-1ra-R nucleic acid molecule, it is typically located at a site 5' from the promoter.

Expression vectors of the invention may be constructed from a starting vector such as a commercially available vector. Such vectors may or may not contain all of the desired flanking sequences. Where one or more of the flanking sequences described herein are not already present in the vector, they may be individually obtained and ligated into the vector. Methods used for obtaining each of the flanking sequences are well known to one skilled in the art.

Preferred vectors for practicing this invention are those which are compatible with bacterial, insect, and mammalian host cells. Such vectors include, *inter alia*, pCRII, pCR3, and pcDNA3.1 (Invitrogen, San Diego, CA), pBSII (Stratagene, La Jolla, CA), pET15 (Novagen, Madison, WI), pGEX (Pharmacia Biotech, Piscataway, NJ), pEGFP-N2 (Clontech, Palo Alto, CA), pETL (BlueBacII, Invitrogen), pDSR-alpha (PCT Pub. No. WO 90/14363) and pFastBacDual (Gibco-BRL, Grand Island, NY).

10

15

20

Additional suitable vectors include, but are not limited to, cosmids, plasmids, or modified viruses, but it will be appreciated that the vector system must be compatible with the selected host cell. Such vectors include, but are not limited to plasmids such as Bluescript® plasmid derivatives (a high copy number ColE1-based phagemid, Stratagene Cloning Systems, La Jolla CA), PCR cloning plasmids designed for cloning Taq-amplified PCR products (*e.g.*, TOPOTM TA Cloning® Kit, PCR2.1® plasmid derivatives, Invitrogen, Carlsbad, CA), and mammalian, yeast or virus vectors such as a baculovirus expression system (pBacPAK plasmid derivatives, Clontech, Palo Alto, CA).

After the vector has been constructed and a nucleic acid molecule encoding an IL-1ra-R polypeptide has been inserted into the proper site of the vector, the completed vector may be inserted into a suitable host cell for amplification and/or polypeptide expression. The transformation of an expression vector for an IL-1ra-R polypeptide into a selected host cell may be accomplished by well known methods including methods such as transfection, infection, calcium chloride, electroporation, microinjection, lipofection, DEAE-dextran method, or other known techniques. The method selected will in part be a function of the type of host cell to be used. These methods and other suitable methods are well known to the skilled artisan, and are set forth, for example, in Sambrook *et al.*, *supra*.

Host cells may be prokaryotic host cells (such as *E. coli*) or eukaryotic host cells (such as a yeast, insect, or vertebrate cell). The host cell, when cultured under appropriate conditions, synthesizes an IL-1ra-R polypeptide which can subsequently be collected from the culture medium (if the host cell secretes it into the medium) or directly from the host cell producing it (if it is not secreted). The selection of an appropriate host cell will depend upon various factors, such as desired expression levels, polypeptide modifications that are desirable or necessary for activity (such as glycosylation or phosphorylation) and ease of folding into a biologically active molecule.

A number of suitable host cells are known in the art and many are available from the American Type Culture Collection (ATCC), Manassas, VA.

30

10

15

20

25

Examples include, but are not limited to, mammalian cells, such as Chinese hamster ovary cells (CHO), CHO DHFR(-) cells (Urlaub et al., 1980, Proc. Natl. Acad. Sci. U.S.A. 97:4216-20), human embryonic kidney (HEK) 293 or 293T cells, or 3T3 cells. The selection of suitable mammalian host cells and methods for transformation, culture, amplification, screening, product production, and purification are known in the art. Other suitable mammalian cell lines, are the monkey COS-1 and COS-7 cell lines, and the CV-1 cell line. Further exemplary mammalian host cells include primate cell lines and rodent cell lines, including transformed cell lines. Normal diploid cells, cell strains derived from in vitro culture of primary tissue, as well as primary explants, are also suitable. Candidate cells may be genotypically deficient in the selection gene, or may contain a dominantly acting selection gene. Other suitable mammalian cell lines include but are not limited to, mouse neuroblastoma N2A cells, HeLa, mouse L-929 cells, 3T3 lines derived from Swiss, Balb-c or NIH mice, BHK or HaK hamster cell lines. Each of these cell lines is known by and available to those skilled in the art of protein expression.

Similarly useful as host cells suitable for the present invention are bacterial cells. For example, the various strains of E. coli (e.g., HB101, DH5 Δ , DH10, and MC1061) are well-known as host cells in the field of biotechnology. Various strains of B. subtilis, $Pseudomonas\ spp.$, other $Bacillus\ spp.$, $Streptomyces\ spp.$, and the like may also be employed in this method.

Many strains of yeast cells known to those skilled in the art are also available as host cells for the expression of the polypeptides of the present invention. Preferred yeast cells include, for example, *Saccharomyces cerivisae* and *Pichia pastoris*.

Additionally, where desired, insect cell systems may be utilized in the methods of the present invention. Such systems are described, for example, in Kitts *et al.*, 1993, *Biotechniques*, 14:810-17; Lucklow, 1993, *Curr. Opin. Biotechnol.* 4:564-72; and Lucklow *et al.*, 1993, *J. Virol.*, 67:4566-79. Preferred insect cells are Sf-9 and Hi5 (Invitrogen).

10

15

20

25

30

One may also use transgenic animals to express glycosylated IL-1ra-R polypeptides. For example, one may use a transgenic milk-producing animal (a cow or goat, for example) and obtain the present glycosylated polypeptide in the animal milk. One may also use plants to produce IL-1ra-R polypeptides, however, in general, the glycosylation occurring in plants is different from that produced in mammalian cells, and may result in a glycosylated product which is not suitable for human therapeutic use.

Polypeptide Production

Host cells comprising an IL-1ra-R polypeptide expression vector may be cultured using standard media well known to the skilled artisan. The media will usually contain all nutrients necessary for the growth and survival of the cells. Suitable media for culturing *E. coli* cells include, for example, Luria Broth (LB) and/or Terrific Broth (TB). Suitable media for culturing eukaryotic cells include Roswell Park Memorial Institute medium 1640 (RPMI 1640), Minimal Essential Medium (MEM) and/or Dulbecco's Modified Eagle Medium (DMEM), all of which may be supplemented with serum and/or growth factors as necessary for the particular cell line being cultured. A suitable medium for insect cultures is Grace's medium supplemented with yeastolate, lactalbumin hydrolysate, and/or fetal calf serum as necessary.

Typically, an antibiotic or other compound useful for selective growth of transfected or transformed cells is added as a supplement to the media. The compound to be used will be dictated by the selectable marker element present on the plasmid with which the host cell was transformed. For example, where the selectable marker element is kanamycin resistance, the compound added to the culture medium will be kanamycin. Other compounds for selective growth include ampicillin, tetracycline, and neomycin.

The amount of an IL-1ra-R polypeptide produced by a host cell can be evaluated using standard methods known in the art. Such methods include, without limitation, Western blot analysis, SDS-polyacrylamide gel electrophoresis, non-denaturing gel electrophoresis, High Performance Liquid

10

15

20

25

30

Chromatography (HPLC) separation, immunoprecipitation, and/or activity assays such as DNA binding gel shift assays.

If an IL-1ra-R polypeptide has been designed to be secreted from the host cells, the majority of polypeptide may be found in the cell culture medium. If however, the IL-1ra-R polypeptide is not secreted from the host cells, it will be present in the cytoplasm and/or the nucleus (for eukaryotic host cells) or in the cytosol (for gram-negative bacteria host cells).

For an IL-1ra-R polypeptide situated in the host cell cytoplasm and/or nucleus (for eukaryotic host cells) or in the cytosol (for bacterial host cells), the intracellular material (including inclusion bodies for gram-negative bacteria) can be extracted from the host cell using any standard technique known to the skilled artisan. For example, the host cells can be lysed to release the contents of the periplasm/cytoplasm by French press, homogenization, and/or sonication followed by centrifugation.

If an IL-1ra-R polypeptide has formed inclusion bodies in the cytosol, the inclusion bodies can often bind to the inner and/or outer cellular membranes and thus will be found primarily in the pellet material after centrifugation. The pellet material can then be treated at pH extremes or with a chaotropic agent such as a detergent, guanidine, guanidine derivatives, urea, or urea derivatives in the presence of a reducing agent such as dithiothreitol at alkaline pH or tris carboxyethyl phosphine at acid pH to release, break apart, and solubilize the inclusion bodies. The solubilized IL-1ra-R polypeptide can then be analyzed using gel electrophoresis, immunoprecipitation, or the like. If it is desired to isolate the IL-1ra-R polypeptide, isolation may be accomplished using standard methods such as those described herein and in Marston *et al.*, 1990, *Meth. Enz.*, 182:264-75.

In some cases, an IL-1ra-R polypeptide may not be biologically active upon isolation. Various methods for "refolding" or converting the polypeptide to its tertiary structure and generating disulfide linkages can be used to restore biological activity. Such methods include exposing the solubilized polypeptide to a pH usually above 7 and in the presence of a particular concentration of a

10

15

20

25

30

chaotrope. The selection of chaotrope is very similar to the choices used for inclusion body solubilization, but usually the chaotrope is used at a lower concentration and is not necessarily the same as chaotropes used for the solubilization. In most cases the refolding/oxidation solution will also contain a reducing agent or the reducing agent plus its oxidized form in a specific ratio to generate a particular redox potential allowing for disulfide shuffling to occur in the formation of the protein's cysteine bridges. Some of the commonly used redox couples include cysteine/cystamine, glutathione (GSH)/dithiobis GSH, 2-2-DTT, and dithiothreitol(DTT)/dithiane cupric chloride, mercaptoethanol(bME)/dithio-b(ME). In many instances, a cosolvent may be used or may be needed to increase the efficiency of the refolding, and the more common reagents used for this purpose include glycerol, polyethylene glycol of various molecular weights, arginine and the like.

If inclusion bodies are not formed to a significant degree upon expression of an IL-1ra-R polypeptide, then the polypeptide will be found primarily in the supernatant after centrifugation of the cell homogenate. The polypeptide may be further isolated from the supernatant using methods such as those described herein.

The purification of an IL-1ra-R polypeptide from solution can be accomplished using a variety of techniques. If the polypeptide has been synthesized such that it contains a tag such as Hexahistidine (IL-1ra-R polypeptide/hexaHis) or other small peptide such as FLAG (Eastman Kodak Co., New Haven, CT) or *myc* (Invitrogen, Carlsbad, CA) at either its carboxyl- or amino-terminus, it may be purified in a one-step process by passing the solution through an affinity column where the column matrix has a high affinity for the tag.

For example, polyhistidine binds with great affinity and specificity to nickel. Thus, an affinity column of nickel (such as the Qiagen[®] nickel columns) can be used for purification of IL-1ra-R polypeptide/polyHis. *See*, *e.g.*, *Current Protocols in Molecular Biology* § 10.11.8 (Ausubel *et al.*, eds., Green Publishers Inc. and Wiley and Sons 1993).

10

15

20

25

30

Additionally, IL-1RA-R polypeptides may be purified through the use of a monoclonal antibody that is capable of specifically recognizing and binding to an IL-1ra-R polypeptide.

Other suitable procedures for purification include, without limitation, affinity chromatography, immunoaffinity chromatography, ion exchange chromatography, molecular sieve chromatography, HPLC, electrophoresis (including native gel electrophoresis) followed by gel elution, and preparative isoelectric focusing ("Isoprime" machine/technique, Hoefer Scientific, San Francisco, CA). In some cases, two or more purification techniques may be combined to achieve increased purity.

IL-1ra-R polypeptides may also be prepared by chemical synthesis methods (such as solid phase peptide synthesis) using techniques known in the art such as those set forth by Merrifield *et al.*, 1963, *J. Am. Chem. Soc.* 85:2149; Houghten *et al.*, 1985, *Proc Natl Acad. Sci. USA* 82:5132; and Stewart and Young, *Solid Phase Peptide Synthesis* (Pierce Chemical Co. 1984). Such polypeptides may be synthesized with or without a methionine on the aminoterminus. Chemically synthesized IL-1ra-R polypeptides may be oxidized using methods set forth in these references to form disulfide bridges. Chemically synthesized IL-1ra-R polypeptides are expected to have comparable biological activity to the corresponding IL-1ra-R polypeptides produced recombinantly or purified from natural sources, and thus may be used interchangeably with a recombinant or natural IL-1ra-R polypeptide.

Another means of obtaining IL-1ra-R polypeptide is via purification from biological samples such as source tissues and/or fluids in which the IL-1ra-R polypeptide is naturally found. Such purification can be conducted using methods for protein purification as described herein. The presence of the IL-1ra-R polypeptide during purification may be monitored, for example, using an antibody prepared against recombinantly produced IL-1ra-R polypeptide or peptide fragments thereof.

A number of additional methods for producing nucleic acids and polypeptides are known in the art, and the methods can be used to produce

10

15

20

25

polypeptides having specificity for IL-1ra-R polypeptide. *See, e.g.*, Roberts *et al.*, 1997, *Proc. Natl. Acad. Sci. U.S.A.* 94:12297-303, which describes the production of fusion proteins between an mRNA and its encoded peptide. *See also*, Roberts, 1999, *Curr. Opin. Chem. Biol.* 3:268-73. Additionally, U.S. Patent No. 5,824,469 describes methods for obtaining oligonucleotides capable of carrying out a specific biological function. The procedure involves generating a heterogeneous pool of oligonucleotides, each having a 5' randomized sequence, a central preselected sequence, and a 3' randomized sequence. The resulting heterogeneous pool is introduced into a population of cells that do not exhibit the desired biological function. Subpopulations of the cells are then screened for those that exhibit a predetermined biological function. From that subpopulation, oligonucleotides capable of carrying out the desired biological function are isolated.

U.S. Patent Nos. 5,763,192; 5,814,476; 5,723,323; and 5,817,483 describe processes for producing peptides or polypeptides. This is done by producing stochastic genes or fragments thereof, and then introducing these genes into host cells which produce one or more proteins encoded by the stochastic genes. The host cells are then screened to identify those clones producing peptides or polypeptides having the desired activity.

Another method for producing peptides or polypeptides is described in PCT/US98/20094 (WO99/15650) filed by Athersys, Inc. Known as "Random Activation of Gene Expression for Gene Discovery" (RAGE-GD), the process involves the activation of endogenous gene expression or over-expression of a gene by *in situ* recombination methods. For example, expression of an endogenous gene is activated or increased by integrating a regulatory sequence into the target cell which is capable of activating expression of the gene by non-homologous or illegitimate recombination. The target DNA is first subjected to radiation, and a genetic promoter inserted. The promoter eventually locates a break at the front of a gene, initiating transcription of the gene. This results in expression of the desired peptide or polypeptide.

It will be appreciated that these methods can also be used to create comprehensive IL-1ra-R polypeptide expression libraries, which can subsequently be used for high throughput phenotypic screening in a variety of assays, such as biochemical assays, cellular assays, and whole organism assays (*e.g.*, plant, mouse, etc.).

Synthesis

5

10

15

20

25

30

It will be appreciated by those skilled in the art that the nucleic acid and polypeptide molecules described herein may be produced by recombinant and other means.

Selective Binding Agents

The term "selective binding agent" refers to a molecule that has specificity for one or more IL-1ra-R polypeptides. Suitable selective binding agents include, but are not limited to, antibodies and derivatives thereof, polypeptides, and small molecules. Suitable selective binding agents may be prepared using methods known in the art. An exemplary IL-1RA-R polypeptide selective binding agent of the present invention is capable of binding a certain portion of the IL-1RA-R polypeptide thereby inhibiting the binding of the polypeptide to an IL-1ra-R polypeptide receptor.

Selective binding agents such as antibodies and antibody fragments that bind IL-1ra-R polypeptides are within the scope of the present invention. The antibodies may be polyclonal including monospecific polyclonal; monoclonal (MAbs); recombinant; chimeric; humanized, such as CDR-grafted; human; single chain; and/or bispecific; as well as fragments; variants; or derivatives thereof. Antibody fragments include those portions of the antibody that bind to an epitope on the IL-1RA-R polypeptide. Examples of such fragments include Fab and F(ab') fragments generated by enzymatic cleavage of full-length antibodies. Other binding fragments include those generated by recombinant DNA techniques, such as the expression of recombinant plasmids containing nucleic acid sequences encoding antibody variable regions.

10

15

20

25

30

Polyclonal antibodies directed toward an IL-1ra-R polypeptide generally are produced in animals (e.g., rabbits or mice) by means of multiple subcutaneous or intraperitoneal injections of IL-1ra-R polypeptide and an adjuvant. It may be useful to conjugate an IL-1ra-R polypeptide to a carrier protein that is immunogenic in the species to be immunized, such as keyhole limpet hemocyanin, serum, albumin, bovine thyroglobulin, or soybean trypsin inhibitor. Also, aggregating agents such as alum are used to enhance the immune response. After immunization, the animals are bled and the serum is assayed for anti-IL-1ra-R antibody titer.

Monoclonal antibodies directed toward IL-1ra-R polypeptides are produced using any method that provides for the production of antibody molecules by continuous cell lines in culture. Examples of suitable methods for preparing monoclonal antibodies include the hybridoma methods of Kohler *et al.*, 1975, *Nature* 256:495-97 and the human B-cell hybridoma method (Kozbor, 1984, *J. Immunol.* 133:3001; Brodeur *et al.*, *Monoclonal Antibody Production Techniques and Applications* 51-63 (Marcel Dekker, Inc., 1987). Also provided by the invention are hybridoma cell lines that produce monoclonal antibodies reactive with IL-1ra-R polypeptides.

Monoclonal antibodies of the invention may be modified for use as therapeutics. One embodiment is a "chimeric" antibody in which a portion of the heavy (H) and/or light (L) chain is identical with or homologous to a corresponding sequence in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is/are identical with or homologous to a corresponding sequence in antibodies derived from another species or belonging to another antibody class or subclass. Also included are fragments of such antibodies, so long as they exhibit the desired biological activity. See U.S. Patent No. 4,816,567; Morrison et al., 1985, Proc. Natl. Acad. Sci. 81:6851-55.

In another embodiment, a monoclonal antibody of the invention is a "humanized" antibody. Methods for humanizing non-human antibodies are well known in the art. See U.S. Patent Nos. 5,585,089 and 5,693,762. Generally, a

15

20

25

30

humanized antibody has one or more amino acid residues introduced into it from a source that is non-human. Humanization can be performed, for example, using methods described in the art (Jones *et al.*, 1986, *Nature* 321:522-25; Riechmann *et al.*, 1998, *Nature* 332:323-27; Verhoeyen *et al.*, 1988, *Science* 239:1534-36), by substituting at least a portion of a rodent complementarity-determining region (CDR) for the corresponding regions of a human antibody.

Also encompassed by the invention are human antibodies that bind IL-1ra-R polypeptides. Using transgenic animals (e.g., mice) that are capable of producing a repertoire of human antibodies in the absence of endogenous immunoglobulin production such antibodies are produced by immunization with an IL-1ra-R polypeptide antigen (i.e., having at least 6 contiguous amino acids), optionally conjugated to a carrier. See, e.g., Jakobovits et al., 1993, Proc. Natl. Acad. Sci. 90:2551-55; Jakobovits et al., 1993, Nature 362:255-58; Bruggermann et al., 1993, Year in Immuno. 7:33. In one method, such transgenic animals are produced by incapacitating the endogenous loci encoding the heavy and light immunoglobulin chains therein, and inserting loci encoding human heavy and light chain proteins into the genome thereof. Partially modified animals, that is those having less than the full complement of modifications, are then cross-bred to obtain an animal having all of the desired immune system modifications. When administered an immunogen, these transgenic animals produce antibodies with human (rather than, e.g., murine) amino acid sequences, including variable regions which are immunospecific for these antigens. See PCT App. Nos. PCT/US96/05928 and PCT/US93/06926. Additional methods are described in PCT/US91/245 5,545,807, PCT Nos. App. U.S. Patent No. PCT/GB89/01207, and in European Patent Nos. 546073B1 and 546073A1. Human antibodies can also be produced by the expression of recombinant DNA in host cells or by expression in hybridoma cells as described herein.

In an alternative embodiment, human antibodies can also be produced from phage-display libraries (Hoogenboom *et al.*, 1991, *J. Mol. Biol.* 227:381; Marks *et al.*, 1991, *J. Mol. Biol.* 222:581). These processes mimic immune selection through the display of antibody repertoires on the surface of filamentous

10

15

20

25

30

bacteriophage, and subsequent selection of phage by their binding to an antigen of choice. One such technique is described in PCT App. No. PCT/US98/17364, which describes the isolation of high affinity and functional agonistic antibodies for MPL- and msk- receptors using such an approach.

Chimeric, CDR grafted, and humanized antibodies are typically produced by recombinant methods. Nucleic acids encoding the antibodies are introduced into host cells and expressed using materials and procedures described herein. In a preferred embodiment, the antibodies are produced in mammalian host cells, such as CHO cells. Monoclonal (e.g., human) antibodies may be produced by the expression of recombinant DNA in host cells or by expression in hybridoma cells as described herein.

The anti-IL-1ra-R antibodies of the invention may be employed in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays (Sola, *Monoclonal Antibodies: A Manual of Techniques* 147-158 (CRC Press, Inc., 1987)) for the detection and quantitation of IL-1ra-R polypeptides. The antibodies will bind IL-1ra-R polypeptides with an affinity that is appropriate for the assay method being employed.

For diagnostic applications, in certain embodiments, anti-IL-1ra-R antibodies may be labeled with a detectable moiety. The detectable moiety can be any one that is capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety may be a radioisotope, such as ³H, ¹⁴C, ³²P, ³⁵S, ¹²⁵I, ⁹⁹Tc, ¹¹¹In, or ⁶⁷Ga; a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin; or an enzyme, such as alkaline phosphatase, E-galactosidase, or horseradish peroxidase (Bayer, *et al.*, 1990, *Meth. Enz.* 184:138-63).

Competitive binding assays rely on the ability of a labeled standard (e.g., an IL-1ra-R polypeptide, or an immunologically reactive portion thereof) to compete with the test sample analyte (an IL-1ra-R polypeptide) for binding with a limited amount of anti-IL-1ra-R antibody. The amount of an IL-1ra-R polypeptide in the test sample is inversely proportional to the amount of standard

10

15

20

25

30

that becomes bound to the antibodies. To facilitate determining the amount of standard that becomes bound, the antibodies typically are insolubilized before or after the competition, so that the standard and analyte that are bound to the antibodies may conveniently be separated from the standard and analyte which remain unbound.

Sandwich assays typically involve the use of two antibodies, each capable of binding to a different immunogenic portion, or epitope, of the protein to be detected and/or quantitated. In a sandwich assay, the test sample analyte is typically bound by a first antibody which is immobilized on a solid support, and thereafter a second antibody binds to the analyte, thus forming an insoluble three-part complex. *See, e.g.*, U.S. Patent No. 4,376,110. The second antibody may itself be labeled with a detectable moiety (direct sandwich assays) or may be measured using an anti-immunoglobulin antibody that is labeled with a detectable moiety (indirect sandwich assays). For example, one type of sandwich assay is an enzyme-linked immunosorbent assay (ELISA), in which case the detectable moiety is an enzyme.

The selective binding agents, including anti-IL-1ra-R antibodies, are also useful for *in vivo* imaging. An antibody labeled with a detectable moiety may be administered to an animal, preferably into the bloodstream, and the presence and location of the labeled antibody in the host assayed. The antibody may be labeled with any moiety that is detectable in an animal, whether by nuclear magnetic resonance, radiology, or other detection means known in the art.

Selective binding agents of the invention, including antibodies, may be used as therapeutics. These therapeutic agents are generally agonists or antagonists, in that they either enhance or reduce, respectively, at least one of the biological activities of an IL-1ra-R polypeptide. In one embodiment, antagonist antibodies of the invention are antibodies or binding fragments thereof which are capable of specifically binding to an IL-1ra-R polypeptide and which are capable of inhibiting or eliminating the functional activity of an IL-1ra-R polypeptide *in vivo* or *in vitro*. In preferred embodiments, the selective binding agent, *e.g.*, an antagonist antibody, will inhibit the functional activity of an IL-1ra-R polypeptide

by at least about 50%, and preferably by at least about 80%. In another embodiment, the selective binding agent may be an anti-IL-1ra-R polypeptide antibody that is capable of interacting with an IL-1ra-R polypeptide binding partner (a ligand or receptor) thereby inhibiting or eliminating IL-1ra-R polypeptide activity *in vitro* or *in vivo*. Selective binding agents, including agonist and antagonist anti-IL-1ra-R polypeptide antibodies, are identified by screening assays that are well known in the art.

The invention also relates to a kit comprising IL-1ra-R selective binding agents (such as antibodies) and other reagents useful for detecting IL-1ra-R polypeptide levels in biological samples. Such reagents may include a detectable label, blocking serum, positive and negative control samples, and detection reagents.

Microarrays

5

10

15

20

25

30

It will be appreciated that DNA microarray technology can be utilized in accordance with the present invention. DNA microarrays are miniature, highdensity arrays of nucleic acids positioned on a solid support, such as glass. Each cell or element within the array contains numerous copies of a single nucleic acid species that acts as a target for hybridization with a complementary nucleic acid In expression profiling using DNA microarray sequence (e.g., mRNA). technology, mRNA is first extracted from a cell or tissue sample and then This material is converted enzymatically to fluorescently labeled cDNA. hybridized to the microarray and unbound cDNA is removed by washing. The expression of discrete genes represented on the array is then visualized by quantitating the amount of labeled cDNA that is specifically bound to each target nucleic acid molecule. In this way, the expression of thousands of genes can be quantitated in a high throughput, parallel manner from a single sample of biological material.

This high throughput expression profiling has a broad range of applications with respect to the IL-1ra-R molecules of the invention, including, but not limited to: the identification and validation of IL-1ra-R disease-related

genes as targets for therapeutics; molecular toxicology of related IL-1ra-R molecules and inhibitors thereof; stratification of populations and generation of surrogate markers for clinical trials; and enhancing related IL-1ra-R polypeptide small molecule drug discovery by aiding in the identification of selective compounds in high throughput screens.

Chemical Derivatives

5

10

15

20

25

Chemically modified derivatives of IL-1ra-R polypeptides may be prepared by one skilled in the art, given the disclosures described herein. IL-1ra-R polypeptide derivatives are modified in a manner that is different – either in the type or location of the molecules naturally attached to the polypeptide. Derivatives may include molecules formed by the deletion of one or more naturally-attached chemical groups. The polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-1ra-R polypeptide, may be modified by the covalent attachment of one or more polymers. For example, the polymer selected is typically water-soluble so that the protein to which it is attached does not precipitate in an aqueous environment, such as a physiological environment. Included within the scope of suitable polymers is a mixture of polymers. Preferably, for therapeutic use of the end-product preparation, the polymer will be pharmaceutically acceptable.

The polymers each may be of any molecular weight and may be branched or unbranched. The polymers each typically have an average molecular weight of between about 2 kDa to about 100 kDa (the term "about" indicating that in preparations of a water-soluble polymer, some molecules will weigh more, some less, than the stated molecular weight). The average molecular weight of each polymer is preferably between about 5 kDa and about 50 kDa, more preferably between about 12 kDa and about 40 kDa and most preferably between about 20 kDa and about 35 kDa.

Suitable water-soluble polymers or mixtures thereof include, but are not limited to, N-linked or O-linked carbohydrates, sugars, phosphates, polyethylene

10

15 `

20

25

30

glycol (PEG) (including the forms of PEG that have been used to derivatize proteins, including mono-(C₁-C₁₀), alkoxy-, or aryloxy-polyethylene glycol), monomethoxy-polyethylene glycol, dextran (such as low molecular weight dextran of, for example, about 6 kD), cellulose, or other carbohydrate based polymers, poly-(N-vinyl pyrrolidone) polyethylene glycol, propylene glycol homopolymers, polypropylene oxide/ethylene oxide co-polymers, polyoxyethylated polyols (e.g., glycerol), and polyvinyl alcohol. Also encompassed by the present invention are bifunctional crosslinking molecules which may be used to prepare covalently attached IL-1ra-R polypeptide multimers.

In general, chemical derivatization may be performed under any suitable condition used to react a protein with an activated polymer molecule. Methods for preparing chemical derivatives of polypeptides will generally comprise the steps of: (a) reacting the polypeptide with the activated polymer molecule (such as a reactive ester or aldehyde derivative of the polymer molecule) under conditions whereby the polypeptide comprising the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or other IL-1ra-R polypeptide, becomes attached to one or more polymer molecules, and (b) obtaining the reaction products. The optimal reaction conditions will be determined based on known parameters and the desired result. For example, the larger the ratio of polymer molecules to protein, the greater the percentage of attached polymer molecule. In one embodiment, the IL-1ra-R polypeptide derivative may have a single polymer molecule moiety at the amino-terminus. See, e.g., U.S. Patent No. 5,234,784.

The pegylation of a polypeptide may be specifically carried out using any of the pegylation reactions known in the art. Such reactions are described, for example, in the following references: Francis *et al.*, 1992, *Focus on Growth Factors* 3:4-10; European Patent Nos. 0154316 and 0401384; and U.S. Patent No. 4,179,337. For example, pegylation may be carried out via an acylation reaction or an alkylation reaction with a reactive polyethylene glycol molecule (or an analogous reactive water-soluble polymer) as described herein. For the acylation

10

15

20

25

Patent No. 5,557,032.

reactions, a selected polymer should have a single reactive ester group. For reductive alkylation, a selected polymer should have a single reactive aldehyde group. A reactive aldehyde is, for example, polyethylene glycol propionaldehyde, which is water stable, or mono C_1 - C_{10} alkoxy or aryloxy derivatives thereof (*see* U.S. Patent No. 5,252,714).

In another embodiment, IL-1ra-R polypeptides may be chemically coupled to biotin. The biotin/IL-1ra-R polypeptide molecules are then allowed to bind to avidin, resulting in tetravalent avidin/biotin/IL-1ra-R polypeptide molecules. IL-1ra-R polypeptides may also be covalently coupled to dinitrophenol (DNP) or trinitrophenol (TNP) and the resulting conjugates precipitated with anti-DNP or anti-TNP-IgM to form decameric conjugates with a valency of 10.

Generally, conditions that may be alleviated or modulated by the administration of the present IL-1ra-R polypeptide derivatives include those described herein for IL-1ra-R polypeptides. However, the IL-1ra-R polypeptide derivatives disclosed herein may have additional activities, enhanced or reduced biological activity, or other characteristics, such as increased or decreased half-life, as compared to the non-derivatized molecules.

Genetically Engineered Non-Human Animals

Additionally included within the scope of the present invention are non-human animals such as mice, rats, or other rodents; rabbits, goats, sheep, or other farm animals, in which the genes encoding native IL-1ra-R polypeptide have been disrupted (i.e., "knocked out") such that the level of expression of IL-1ra-R polypeptide is significantly decreased or completely abolished. Such animals may be prepared using techniques and methods such as those described in U.S.

The present invention further includes non-human animals such as mice, rats, or other rodents; rabbits, goats, sheep, or other farm animals, in which either the native form of an IL-1ra-R gene for that animal or a heterologous IL-1ra-R

gene is over-expressed by the animal, thereby creating a "transgenic" animal.

10

15

20

25

30

Such transgenic animals may be prepared using well known methods such as those described in U.S. Patent No 5,489,743 and PCT Pub. No. WO 94/28122.

The present invention further includes non-human animals in which the promoter for one or more of the IL-1ra-R polypeptides of the present invention is either activated or inactivated (e.g., by using homologous recombination methods) to alter the level of expression of one or more of the native IL-1ra-R polypeptides.

These non-human animals may be used for drug candidate screening. In such screening, the impact of a drug candidate on the animal may be measured. For example, drug candidates may decrease or increase the expression of the IL-1ra-R gene. In certain embodiments, the amount of IL-1ra-R polypeptide that is produced may be measured after the exposure of the animal to the drug candidate. Additionally, in certain embodiments, one may detect the actual impact of the drug candidate on the animal. For example, over-expression of a particular gene may result in, or be associated with, a disease or pathological condition. In such cases, one may test a drug candidate's ability to decrease expression of the gene or its ability to prevent or inhibit a pathological condition. In other examples, the production of a particular metabolic product such as a fragment of a polypeptide, may result in, or be associated with, a disease or pathological condition. In such cases, one may test a drug candidate's ability to decrease the production of such a metabolic product or its ability to prevent or inhibit a pathological condition.

Assaying for Other Modulators of IL-1ra-R Polypeptide Activity

In some situations, it may be desirable to identify molecules that are modulators, *i.e.*, agonists or antagonists, of the activity of IL-1ra-R polypeptide. Natural or synthetic molecules that modulate IL-1ra-R polypeptide may be identified using one or more screening assays, such as those described herein. Such molecules may be administered either in an *ex vivo* manner or in an *in vivo* manner by injection, or by oral delivery, implantation device, or the like.

"Test molecule" refers to a molecule that is under evaluation for the ability to modulate (*i.e.*, increase or decrease) the activity of an IL-1ra-R polypeptide. Most commonly, a test molecule will interact directly with an IL-1ra-R

10

15

20

25

polypeptide. However, it is also contemplated that a test molecule may also modulate IL-1ra-R polypeptide activity indirectly, such as by affecting IL-1ra-R gene expression, or by binding to an IL-1ra-R polypeptide binding partner (*e.g.*, receptor or ligand). In one embodiment, a test molecule will bind to an IL-1ra-R polypeptide with an affinity constant of at least about 10⁻⁶ M, preferably about 10⁻⁸ M, more preferably about 10⁻⁹ M, and even more preferably about 10⁻¹⁰ M.

Methods for identifying compounds that interact with IL-1ra-R polypeptides are encompassed by the present invention. In certain embodiments, an IL-1ra-R polypeptide is incubated with a test molecule under conditions that permit the interaction of the test molecule with an IL-1ra-R polypeptide, and the extent of the interaction is measured. The test molecule can be screened in a substantially purified form or in a crude mixture.

In certain embodiments, an IL-1ra-R polypeptide agonist or antagonist may be a protein, peptide, carbohydrate, lipid, or small molecular weight molecule that interacts with IL-1ra-R polypeptide to regulate its activity. Molecules which regulate IL-1ra-R polypeptide expression include nucleic acids which are complementary to nucleic acids encoding an IL-1ra-R polypeptide, or are complementary to nucleic acids sequences which direct or control the expression of IL-1ra-R polypeptide, and which act as anti-sense regulators of expression.

Once a test molecule has been identified as interacting with an IL-1ra-R polypeptide, the molecule may be further evaluated for its ability to increase or decrease IL-1ra-R polypeptide activity. The measurement of the interaction of a test molecule with IL-1ra-R polypeptide may be carried out in several formats, including cell-based binding assays, membrane binding assays, solution-phase assays, and immunoassays. In general, a test molecule is incubated with an IL-1ra-R polypeptide for a specified period of time, and IL-1ra-R polypeptide activity is determined by one or more assays for measuring biological activity.

The interaction of test molecules with IL-1ra-R polypeptides may also be assayed directly using polyclonal or monoclonal antibodies in an immunoassay.

10

15

20

25

30

Alternatively, modified forms of IL-1ra-R polypeptides containing epitope tags as described herein may be used in solution and immunoassays.

In the event that IL-1ra-R polypeptides display biological activity through an interaction with a binding partner (e.g., a receptor or a ligand), a variety of in vitro assays may be used to measure the binding of an IL-1ra-R polypeptide to the corresponding binding partner (such as a selective binding agent, receptor, or ligand). These assays may be used to screen test molecules for their ability to increase or decrease the rate and/or the extent of binding of an IL-1ra-R polypeptide to its binding partner. In one assay, an IL-1ra-R polypeptide is immobilized in the wells of a microtiter plate. Radiolabeled IL-1ra-R polypeptide binding partner (for example, iodinated IL-1ra-R polypeptide binding partner) and a test molecule can then be added either one at a time (in either order) or simultaneously to the wells. After incubation, the wells can be washed and counted for radioactivity, using a scintillation counter, to determine the extent to which the binding partner bound to the IL-1ra-R polypeptide. Typically, a molecule will be tested over a range of concentrations, and a series of control wells lacking one or more elements of the test assays can be used for accuracy in the evaluation of the results. An alternative to this method involves reversing the "positions" of the proteins, i.e., immobilizing IL-1ra-R polypeptide binding partner to the microtiter plate wells, incubating with the test molecule and radiolabeled IL-1ra-R polypeptide, and determining the extent of IL-1ra-R polypeptide binding. See, e.g., Current Protocols in Molecular Biology, chap. 18 (Ausubel et al., eds., Green Publishers Inc. and Wiley and Sons 1995).

As an alternative to radiolabeling, an IL-1ra-R polypeptide or its binding partner may be conjugated to biotin, and the presence of biotinylated protein can then be detected using streptavidin linked to an enzyme, such as horse radish peroxidase (HRP) or alkaline phosphatase (AP), which can be detected colorometrically, or by fluorescent tagging of streptavidin. An antibody directed to an IL-1ra-R polypeptide or to an IL-1ra-R polypeptide binding partner, and which is conjugated to biotin, may also be used for purposes of detection

10

15

20

following incubation of the complex with enzyme-linked streptavidin linked to AP or HRP.

A IL-1ra-R polypeptide or an IL-1ra-R polypeptide binding partner can also be immobilized by attachment to agarose beads, acrylic beads, or other types of such inert solid phase substrates. The substrate-protein complex can be placed in a solution containing the complementary protein and the test compound. After incubation, the beads can be precipitated by centrifugation, and the amount of binding between an IL-1ra-R polypeptide and its binding partner can be assessed using the methods described herein. Alternatively, the substrate-protein complex can be immobilized in a column with the test molecule and complementary protein passing through the column. The formation of a complex between an IL-1ra-R polypeptide and its binding partner can then be assessed using any of the techniques described herein (e.g., radiolabelling or antibody binding).

Another *in vitro* assay that is useful for identifying a test molecule which increases or decreases the formation of a complex between an IL-1ra-R polypeptide binding protein and an IL-1ra-R polypeptide binding partner is a surface plasmon resonance detector system such as the BIAcore assay system (Pharmacia, Piscataway, NJ). The BIAcore system is utilized as specified by the manufacturer. This assay essentially involves the covalent binding of either IL-1ra-R polypeptide or an IL-1ra-R polypeptide binding partner to a dextran-coated sensor chip that is located in a detector. The test compound and the other complementary protein can then be injected, either simultaneously or sequentially, into the chamber containing the sensor chip. The amount of complementary protein that binds can be assessed based on the change in molecular mass that is physically associated with the dextran-coated side of the sensor chip, with the change in molecular mass being measured by the detector system.

In some cases, it may be desirable to evaluate two or more test compounds together for their ability to increase or decrease the formation of a complex between an IL-1ra-R polypeptide and an IL-1ra-R polypeptide binding partner. In these cases, the assays set forth herein can be readily modified by adding such

10

15

20

25

additional test compound(s) either simultaneously with, or subsequent to, the first test compound. The remainder of the steps in the assay are as set forth herein.

In vitro assays such as those described herein may be used advantageously to screen large numbers of compounds for an effect on the formation of a complex between an IL-1ra-R polypeptide and IL-1ra-R polypeptide binding partner. The assays may be automated to screen compounds generated in phage display, synthetic peptide, and chemical synthesis libraries.

Compounds which increase or decrease the formation of a complex between an IL-1ra-R polypeptide and an IL-1ra-R polypeptide binding partner may also be screened in cell culture using cells and cell lines expressing either IL-1ra-R polypeptide or IL-1ra-R polypeptide binding partner. Cells and cell lines may be obtained from any mammal, but preferably will be from human or other primate, canine, or rodent sources. The binding of an IL-1ra-R polypeptide to cells expressing IL-1ra-R polypeptide binding partner at the surface is evaluated in the presence or absence of test molecules, and the extent of binding may be determined by, for example, flow cytometry using a biotinylated antibody to an IL-1ra-R polypeptide binding partner. Cell culture assays can be used advantageously to further evaluate compounds that score positive in protein binding assays described herein.

Cell cultures can also be used to screen the impact of a drug candidate. For example, drug candidates may decrease or increase the expression of the IL-1ra-R gene. In certain embodiments, the amount of IL-1ra-R polypeptide or an IL-1ra-R polypeptide fragment that is produced may be measured after exposure of the cell culture to the drug candidate. In certain embodiments, one may detect the actual impact of the drug candidate on the cell culture. For example, the over-expression of a particular gene may have a particular impact on the cell culture. In such cases, one may test a drug candidate's ability to increase or decrease the expression of the gene or its ability to prevent or inhibit a particular impact on the cell culture. In other examples, the production of a particular metabolic product such as a fragment of a polypeptide, may result in, or be associated with, a disease

or pathological condition. In such cases, one may test a drug candidate's ability to decrease the production of such a metabolic product in a cell culture.

Internalizing Proteins

5

10

15

20

25

The tat protein sequence (from HIV) can be used to internalize proteins into a cell. See, e.g., Falwell et al., 1994, Proc. Natl. Acad. Sci. U.S.A. 91:664-68. For example, an 11 amino acid sequence (Y-G-R-K-K-R-R-Q-R-R-R; SEQ ID NO: 18) of the HIV tat protein (termed the "protein transduction domain," or TAT PDT) has been described as mediating delivery across the cytoplasmic membrane and the nuclear membrane of a cell. See Schwarze et al., 1999, Science 285:1569-72; and Nagahara et al., 1998, Nat. Med. 4:1449-52. In these procedures, FITCconstructs (FITC-labeled G-G-G-Y-G-R-K-K-R-R-Q-R-R-R; SEQ ID NO: 19), which penetrate tissues following intraperitoneal administration, are prepared, and the binding of such constructs to cells is detected by fluorescence-activated cell sorting (FACS) analysis. Cells treated with a tat-E-gal fusion protein will demonstrate E-gal activity. Following injection, expression of such a construct can be detected in a number of tissues, including liver, kidney, lung, heart, and brain tissue. It is believed that such constructs undergo some degree of unfolding in order to enter the cell, and as such, may require a refolding following entry into the cell.

It will thus be appreciated that the *tat* protein sequence may be used to internalize a desired polypeptide into a cell. For example, using the *tat* protein sequence, an IL-1ra-R antagonist (such as an anti-IL-1ra-R selective binding agent, small molecule, soluble receptor, or antisense oligonucleotide) can be administered intracellularly to inhibit the activity of an IL-1ra-R molecule. As used herein, the term "IL-1ra-R molecule" refers to both IL-1ra-R nucleic acid molecules and IL-1ra-R polypeptides as defined herein. Where desired, the IL-1ra-R protein itself may also be internally administered to a cell using these procedures. *See also*, Straus, 1999, *Science* 285:1466-67.

In accordance with certain embodiments of the invention, it may be useful to be able to determine the source of a certain cell type associated with an IL-1ra-R polypeptide. For example, it may be useful to determine the origin of a disease or pathological condition as an aid in selecting an appropriate therapy. In certain embodiments, nucleic acids encoding an IL-1ra-R polypeptide can be used as a probe to identify cells described herein by screening the nucleic acids of the cells with such a probe. In other embodiments, one may use anti-IL-1ra-R polypeptide antibodies to test for the presence of IL-1ra-R polypeptide in cells, and thus, determine if such cells are of the types described herein.

10

15

20

25

30

5

IL-1ra-R Polypeptide Compositions and Administration

Therapeutic compositions are within the scope of the present invention. Such IL-1RA-R polypeptide pharmaceutical compositions may comprise a therapeutically effective amount of an IL-1ra-R polypeptide or an IL-1ra-R nucleic acid molecule in admixture with a pharmaceutically or physiologically acceptable formulation agent selected for suitability with the mode of administration. Pharmaceutical compositions may comprise a therapeutically effective amount of one or more IL-1ra-R polypeptide selective binding agents in admixture with a pharmaceutically or physiologically acceptable formulation agent selected for suitability with the mode of administration.

Acceptable formulation materials preferably are nontoxic to recipients at the dosages and concentrations employed.

The pharmaceutical composition may contain formulation materials for modifying, maintaining, or preserving, for example, the pH, osmolarity, viscosity, clarity, color, isotonicity, odor, sterility, stability, rate of dissolution or release, adsorption, or penetration of the composition. Suitable formulation materials include, but are not limited to, amino acids (such as glycine, glutamine, asparagine, arginine, or lysine), antimicrobials, antioxidants (such as ascorbic acid, sodium sulfite, or sodium hydrogen-sulfite), buffers (such as borate, bicarbonate, Tris-HCl, citrates, phosphates, or other organic acids), bulking agents (such as mannitol or glycine), chelating agents (such as ethylenediamine

10

15

20

25

30

(such caffeine. complexing agents as (EDTA)), tetraacetic acid polyvinylpyrrolidone, beta-cyclodextrin, or hydroxypropyl-beta-cyclodextrin), fillers, monosaccharides, disaccharides, and other carbohydrates (such as glucose, mannose, or dextrins), proteins (such as serum albumin, gelatin, or immunoglobulins), coloring, flavoring and diluting agents, emulsifying agents, hydrophilic polymers (such as polyvinylpyrrolidone), low molecular weight polypeptides, salt-forming counterions (such as sodium), preservatives (such as benzalkonium chloride, benzoic acid, salicylic acid, thimerosal, phenethyl alcohol, methylparaben, propylparaben, chlorhexidine, sorbic acid, or hydrogen peroxide), solvents (such as glycerin, propylene glycol, or polyethylene glycol), sugar alcohols (such as mannitol or sorbitol), suspending agents, surfactants or wetting agents (such as pluronics; PEG; sorbitan esters; polysorbates such as polysorbate 20 or polysorbate 80; triton; tromethamine; lecithin; cholesterol or tyloxapal), stability enhancing agents (such as sucrose or sorbitol), tonicity enhancing agents (such as alkali metal halides - preferably sodium or potassium chloride - or mannitol sorbitol), delivery vehicles, diluents, excipients and/or pharmaceutical adjuvants. See Remington's Pharmaceutical Sciences (18th Ed., A.R. Gennaro, ed., Mack Publishing Company 1990.

The optimal pharmaceutical composition will be determined by a skilled artisan depending upon, for example, the intended route of administration, delivery format, and desired dosage. See, e.g., Remington's Pharmaceutical Sciences, supra. Such compositions may influence the physical state, stability, rate of in vivo release, and rate of in vivo clearance of the IL-1ra-R molecule.

The primary vehicle or carrier in a pharmaceutical composition may be either aqueous or non-aqueous in nature. For example, a suitable vehicle or carrier for injection may be water, physiological saline solution, or artificial cerebrospinal fluid, possibly supplemented with other materials common in compositions for parenteral administration. Neutral buffered saline or saline mixed with serum albumin are further exemplary vehicles. Other exemplary pharmaceutical compositions comprise Tris buffer of about pH 7.0-8.5, or acetate buffer of about pH 4.0-5.5, which may further include sorbitol or a suitable

10

15

20

25

substitute. In one embodiment of the present invention, IL-1ra-R polypeptide compositions may be prepared for storage by mixing the selected composition having the desired degree of purity with optional formulation agents (*Remington's Pharmaceutical Sciences*, *supra*) in the form of a lyophilized cake or an aqueous solution. Further, the IL-1ra-R polypeptide product may be formulated as a lyophilizate using appropriate excipients such as sucrose.

The IL-1ra-R polypeptide pharmaceutical compositions can be selected for parenteral delivery. Alternatively, the compositions may be selected for inhalation or for delivery through the digestive tract, such as orally. The preparation of such pharmaceutically acceptable compositions is within the skill of the art.

The formulation components are present in concentrations that are acceptable to the site of administration. For example, buffers are used to maintain the composition at physiological pH or at a slightly lower pH, typically within a pH range of from about 5 to about 8.

When parenteral administration is contemplated, the therapeutic compositions for use in this invention may be in the form of a pyrogen-free, parenterally acceptable, aqueous solution comprising the desired IL-1ra-R molecule in a pharmaceutically acceptable vehicle. A particularly suitable vehicle for parenteral injection is sterile distilled water in which an IL-1ra-R molecule is formulated as a sterile, isotonic solution, properly preserved. Yet another preparation can involve the formulation of the desired molecule with an agent, such as injectable microspheres, bio-erodible particles, polymeric compounds (such as polylactic acid or polyglycolic acid), beads, or liposomes, that provides for the controlled or sustained release of the product which may then be delivered via a depot injection. Hyaluronic acid may also be used, and this may have the effect of promoting sustained duration in the circulation. Other suitable means for the introduction of the desired molecule include implantable drug delivery devices.

In one embodiment, a pharmaceutical composition may be formulated for inhalation. For example, IL-1ra-R polypeptide may be formulated as a dry

10

15

20

25

30

powder for inhalation. IL-1ra-R polypeptide or nucleic acid molecule inhalation solutions may also be formulated with a propellant for aerosol delivery. In yet another embodiment, solutions may be nebulized. Pulmonary administration is further described in PCT Pub. No. WO 94/20069, which describes the pulmonary delivery of chemically modified proteins.

It is also contemplated that certain formulations may be administered orally. In one embodiment of the present invention, IL-1ra-R polypeptides that are administered in this fashion can be formulated with or without those carriers customarily used in the compounding of solid dosage forms such as tablets and capsules. For example, a capsule may be designed to release the active portion of the formulation at the point in the gastrointestinal tract when bioavailability is maximized and pre-systemic degradation is minimized. Additional agents can be included to facilitate absorption of the IL-1ra-R polypeptide. Diluents, flavorings, low melting point waxes, vegetable oils, lubricants, suspending agents, tablet disintegrating agents, and binders may also be employed.

Another pharmaceutical composition may involve an effective quantity of IL-1ra-R polypeptides in a mixture with non-toxic excipients that are suitable for the manufacture of tablets. By dissolving the tablets in sterile water, or another appropriate vehicle, solutions can be prepared in unit-dose form. Suitable excipients include, but are not limited to, inert diluents, such as calcium carbonate, sodium carbonate or bicarbonate, lactose, or calcium phosphate; or binding agents, such as starch, gelatin, or acacia; or lubricating agents such as magnesium stearate, stearic acid, or talc.

Additional IL-1ra-R polypeptide pharmaceutical compositions will be evident to those skilled in the art, including formulations involving IL-1ra-R polypeptides in sustained- or controlled-delivery formulations. Techniques for formulating a variety of other sustained- or controlled-delivery means, such as liposome carriers, bio-erodible microparticles or porous beads and depot injections, are also known to those skilled in the art. *See*, *e.g.*, PCT/US93/00829, which describes the controlled release of porous polymeric microparticles for the delivery of pharmaceutical compositions.

10

15

20

25

30

Additional sustained-release include examples of preparations semipermeable polymer matrices in the form of shaped articles, e.g. films, or microcapsules. Sustained release matrices may include polyesters, hydrogels, polylactides (U.S. Patent No. 3,773,919 and European Patent No. 058481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., 1983, Biopolymers 22:547-56), poly(2-hydroxyethyl-methacrylate) (Langer et al., 1981, J. Biomed. Mater. Res. 15:167-277 and Langer, 1982, Chem. Tech. 12:98-105), ethylene vinyl acetate (Langer et al., supra) or poly-D(-)-3-hydroxybutyric acid (European Patent No. 133988). Sustained-release compositions may also include liposomes, which can be prepared by any of several methods known in the art. See, e.g., Eppstein et al., 1985, Proc. Natl. Acad. Sci. USA 82:3688-92; and European Patent Nos. 036676, 088046, and 143949.

The IL-1ra-R pharmaceutical composition to be used for *in vivo* administration typically must be sterile. This may be accomplished by filtration through sterile filtration membranes. Where the composition is lyophilized, sterilization using this method may be conducted either prior to, or following, lyophilization and reconstitution. The composition for parenteral administration may be stored in lyophilized form or in a solution. In addition, parenteral compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

Once the pharmaceutical composition has been formulated, it may be stored in sterile vials as a solution, suspension, gel, emulsion, solid, or as a dehydrated or lyophilized powder. Such formulations may be stored either in a ready-to-use form or in a form (e.g., lyophilized) requiring reconstitution prior to administration.

In a specific embodiment, the present invention is directed to kits for producing a single-dose administration unit. The kits may each contain both a first container having a dried protein and a second container having an aqueous formulation. Also included within the scope of this invention are kits containing

10

15

20

25

30

single and multi-chambered pre-filled syringes (e.g., liquid syringes and lyosyringes).

The effective amount of an IL-1ra-R pharmaceutical composition to be employed therapeutically will depend, for example, upon the therapeutic context and objectives. One skilled in the art will appreciate that the appropriate dosage levels for treatment will thus vary depending, in part, upon the molecule delivered, the indication for which the IL-1ra-R molecule is being used, the route of administration, and the size (body weight, body surface, or organ size) and condition (the age and general health) of the patient. Accordingly, the clinician may titer the dosage and modify the route of administration to obtain the optimal therapeutic effect. A typical dosage may range from about 0.1 Tg/kg to up to about 100 mg/kg or more, depending on the factors mentioned above. In other embodiments, the dosage may range from 0.1 Tg/kg up to about 100 mg/kg; or 1 Tg/kg up to about 100 mg/kg; or 5 Tg/kg up to about 100 mg/kg.

The frequency of dosing will depend upon the pharmacokinetic parameters of the IL-1ra-R molecule in the formulation being used. Typically, a clinician will administer the composition until a dosage is reached that achieves the desired effect. The composition may therefore be administered as a single dose, as two or more doses (which may or may not contain the same amount of the desired molecule) over time, or as a continuous infusion via an implantation device or catheter. Further refinement of the appropriate dosage is routinely made by those of ordinary skill in the art and is within the ambit of tasks routinely performed by them. Appropriate dosages may be ascertained through use of appropriate doseresponse data.

The route of administration of the pharmaceutical composition is in accord with known methods, *e.g.*, orally; through injection by intravenous, intraperitoneal, intracerebral (intraparenchymal), intracerebroventricular, intramuscular, intraocular, intraarterial, intraportal, or intralesional routes; by sustained release systems; or by implantation devices. Where desired, the compositions may be administered by bolus injection or continuously by infusion, or by implantation device.

10

15

20

25

30

Alternatively or additionally, the composition may be administered locally via implantation of a membrane, sponge, or other appropriate material onto which the desired molecule has been absorbed or encapsulated. Where an implantation device is used, the device may be implanted into any suitable tissue or organ, and delivery of the desired molecule may be via diffusion, timed-release bolus, or continuous administration.

In some cases, it may be desirable to use IL-1ra-R polypeptide pharmaceutical compositions in an *ex vivo* manner. In such instances, cells, tissues, or organs that have been removed from the patient are exposed to IL-1ra-R polypeptide pharmaceutical compositions after which the cells, tissues, or organs are subsequently implanted back into the patient.

In other cases, an IL-1ra-R polypeptide can be delivered by implanting certain cells that have been genetically engineered, using methods such as those described herein, to express and secrete the IL-1ra-R polypeptide. Such cells may be animal or human cells, and may be autologous, heterologous, or xenogeneic. Optionally, the cells may be immortalized. In order to decrease the chance of an immunological response, the cells may be encapsulated to avoid infiltration of surrounding tissues. The encapsulation materials are typically biocompatible, semi-permeable polymeric enclosures or membranes that allow the release of the protein product(s) but prevent the destruction of the cells by the patient's immune system or by other detrimental factors from the surrounding tissues.

As discussed herein, it may be desirable to treat isolated cell populations (such as stem cells, lymphocytes, red blood cells, chondrocytes, neurons, and the like) with one or more IL-1ra-R polypeptides. This can be accomplished by exposing the isolated cells to the polypeptide directly, where it is in a form that is permeable to the cell membrane.

Additional embodiments of the present invention relate to cells and methods (e.g., homologous recombination and/or other recombinant production methods) for both the *in vitro* production of therapeutic polypeptides and for the production and delivery of therapeutic polypeptides by gene therapy or cell therapy. Homologous and other recombination methods may be used to modify a

10

15

20

25

30

cell that contains a normally transcriptionally-silent IL-1ra-R gene, or an underexpressed gene, and thereby produce a cell which expresses therapeutically efficacious amounts of IL-1ra-R polypeptides.

Homologous recombination is a technique originally developed for targeting genes to induce or correct mutations in transcriptionally active genes. Kucherlapati, 1989, *Prog. in Nucl. Acid Res. & Mol. Biol.* 36:301. The basic technique was developed as a method for introducing specific mutations into specific regions of the mammalian genome (Thomas *et al.*, 1986, *Cell* 44:419-28; Thomas and Capecchi, 1987, *Cell* 51:503-12; Doetschman *et al.*, 1988, *Proc. Natl. Acad. Sci. U.S.A.* 85:8583-87) or to correct specific mutations within defective genes (Doetschman *et al.*, 1987, *Nature* 330:576-78). Exemplary homologous recombination techniques are described in U.S. Patent No. 5,272,071; European Patent Nos. 9193051 and 505500; PCT/US90/07642, and PCT Pub No. WO 91/09955).

Through homologous recombination, the DNA sequence to be inserted into the genome can be directed to a specific region of the gene of interest by attaching it to targeting DNA. The targeting DNA is a nucleotide sequence that is complementary (homologous) to a region of the genomic DNA. Small pieces of targeting DNA that are complementary to a specific region of the genome are put in contact with the parental strand during the DNA replication process. It is a general property of DNA that has been inserted into a cell to hybridize, and therefore, recombine with other pieces of endogenous DNA through shared homologous regions. If this complementary strand is attached to an oligonucleotide that contains a mutation or a different sequence or an additional nucleotide, it too is incorporated into the newly synthesized strand as a result of the recombination. As a result of the proofreading function, it is possible for the new sequence of DNA to serve as the template. Thus, the transferred DNA is incorporated into the genome.

Attached to these pieces of targeting DNA are regions of DNA that may interact with or control the expression of an IL-1ra-R polypeptide, e.g., flanking sequences. For example, a promoter/enhancer element, a suppressor, or an

exogenous transcription modulatory element is inserted in the genome of the intended host cell in proximity and orientation sufficient to influence the transcription of DNA encoding the desired IL-1ra-R polypeptide. The control element controls a portion of the DNA present in the host cell genome. Thus, the expression of the desired IL-1ra-R polypeptide may be achieved not by transfection of DNA that encodes the IL-1ra-R gene itself, but rather by the use of targeting DNA (containing regions of homology with the endogenous gene of interest) coupled with DNA regulatory segments that provide the endogenous gene sequence with recognizable signals for transcription of an IL-1ra-R gene.

10

15

5

In an exemplary method, the expression of a desired targeted gene in a cell (*i.e.*, a desired endogenous cellular gene) is altered via homologous recombination into the cellular genome at a preselected site, by the introduction of DNA which includes at least a regulatory sequence, an exon, and a splice donor site. These components are introduced into the chromosomal (genomic) DNA in such a manner that this, in effect, results in the production of a new transcription unit (in which the regulatory sequence, the exon, and the splice donor site present in the DNA construct are operatively linked to the endogenous gene). As a result of the introduction of these components into the chromosomal DNA, the expression of the desired endogenous gene is altered.

20

Altered gene expression, as described herein, encompasses activating (or causing to be expressed) a gene which is normally silent (unexpressed) in the cell as obtained, as well as increasing the expression of a gene which is not expressed at physiologically significant levels in the cell as obtained. The embodiments further encompass changing the pattern of regulation or induction such that it is different from the pattern of regulation or induction that occurs in the cell as obtained, and reducing (including eliminating) the expression of a gene which is expressed in the cell as obtained.

30

25

One method by which homologous recombination can be used to increase, or cause, IL-1ra-R polypeptide production from a cell's endogenous IL-1ra-R gene involves first using homologous recombination to place a recombination sequence from a site-specific recombination system (e.g., Cre/loxP, FLP/FRT)

10

15

20

25

30

(Sauer, 1994, Curr. Opin. Biotechnol., 5:521-27; Sauer, 1993, Methods Enzymol., 225:890-900) upstream of (i.e., 5' to) the cell's endogenous genomic IL-1ra-R polypeptide coding region. A plasmid containing a recombination site homologous to the site that was placed just upstream of the genomic IL-1ra-R polypeptide coding region is introduced into the modified cell line along with the appropriate recombinase enzyme. This recombinase causes the plasmid to integrate, via the plasmid's recombination site, into the recombination site located just upstream of the genomic IL-1ra-R polypeptide coding region in the cell line (Baubonis and Sauer, 1993, Nucleic Acids Res. 21:2025-29; O'Gorman et al., 1991, Science 251:1351-55). Any flanking sequences known to increase transcription (e.g., enhancer/promoter, intron, translational enhancer), if properly positioned in this plasmid, would integrate in such a manner as to create a new or modified transcriptional unit resulting in de novo or increased IL-1ra-R polypeptide production from the cell's endogenous IL-1ra-R gene.

A further method to use the cell line in which the site specific recombination sequence had been placed just upstream of the cell's endogenous genomic IL-1ra-R polypeptide coding region is to use homologous recombination to introduce a second recombination site elsewhere in the cell line's genome. The appropriate recombinase enzyme is then introduced into the two-recombination-site cell line, causing a recombination event (deletion, inversion, and translocation) (Sauer, 1994, *Curr. Opin. Biotechnol.*, 5:521-27; Sauer, 1993, *Methods Enzymol.*, 225:890-900) that would create a new or modified transcriptional unit resulting in *de novo* or increased IL-1ra-R polypeptide production from the cell's endogenous IL-1ra-R gene.

An additional approach for increasing, or causing, the expression of IL-1ra-R polypeptide from a cell's endogenous IL-1ra-R gene involves increasing, or causing, the expression of a gene or genes (e.g., transcription factors) and/or decreasing the expression of a gene or genes (e.g., transcriptional repressors) in a manner which results in *de novo* or increased IL-1ra-R polypeptide production from the cell's endogenous IL-1ra-R gene. This method includes the introduction of a non-naturally occurring polypeptide (e.g., a polypeptide comprising a site

10

15

20

25

30

specific DNA binding domain fused to a transcriptional factor domain) into the cell such that *de novo* or increased IL-1ra-R polypeptide production from the cell's endogenous IL-1ra-R gene results.

The present invention further relates to DNA constructs useful in the method of altering expression of a target gene. In certain embodiments, the exemplary DNA constructs comprise: (a) one or more targeting sequences, (b) a regulatory sequence, (c) an exon, and (d) an unpaired splice-donor site. The targeting sequence in the DNA construct directs the integration of elements (a) - (d) into a target gene in a cell such that the elements (b) - (d) are operatively linked to sequences of the endogenous target gene. In another embodiment, the DNA constructs comprise: (a) one or more targeting sequences, (b) a regulatory sequence, (c) an exon, (d) a splice-donor site, (e) an intron, and (f) a splice-acceptor site, wherein the targeting sequence directs the integration of elements (a) - (f) such that the elements of (b) - (f) are operatively linked to the endogenous gene. The targeting sequence is homologous to the preselected site in the cellular chromosomal DNA with which homologous recombination is to occur. In the construct, the exon is generally 3' of the regulatory sequence and the splice-donor site is 3' of the exon.

If the sequence of a particular gene is known, such as the nucleic acid sequence of IL-1ra-R polypeptide presented herein, a piece of DNA that is complementary to a selected region of the gene can be synthesized or otherwise obtained, such as by appropriate restriction of the native DNA at specific recognition sites bounding the region of interest. This piece serves as a targeting sequence upon insertion into the cell and will hybridize to its homologous region within the genome. If this hybridization occurs during DNA replication, this piece of DNA, and any additional sequence attached thereto, will act as an Okazaki fragment and will be incorporated into the newly synthesized daughter strand of DNA. The present invention, therefore, includes nucleotides encoding an IL-1ra-R polypeptide, which nucleotides may be used as targeting sequences.

IL-1ra-R polypeptide cell therapy, e.g., the implantation of cells producing IL-1ra-R polypeptides, is also contemplated. This embodiment involves

10

15

20

25

30

implanting cells capable of synthesizing and secreting a biologically active form of IL-1ra-R polypeptide. Such IL-1ra-R polypeptide-producing cells can be cells that are natural producers of IL-1ra-R polypeptides or may be recombinant cells whose ability to produce IL-1ra-R polypeptides has been augmented by transformation with a gene encoding the desired IL-1ra-R polypeptide or with a gene augmenting the expression of IL-1ra-R polypeptide. Such a modification may be accomplished by means of a vector suitable for delivering the gene as well as promoting its expression and secretion. In order to minimize a potential immunological reaction in patients being administered an IL-1ra-R polypeptide, as may occur with the administration of a polypeptide of a foreign species, it is preferred that the natural cells producing IL-1ra-R polypeptide be of human origin and produce human IL-1ra-R polypeptide. Likewise, it is preferred that the recombinant cells producing IL-1ra-R polypeptide be transformed with an expression vector containing a gene encoding a human IL-1ra-R polypeptide.

Implanted cells may be encapsulated to avoid the infiltration of surrounding tissue. Human or non-human animal cells may be implanted in patients in biocompatible, semipermeable polymeric enclosures or membranes that allow the release of IL-1ra-R polypeptide, but that prevent the destruction of the cells by the patient's immune system or by other detrimental factors from the surrounding tissue. Alternatively, the patient's own cells, transformed to produce IL-1ra-R polypeptides *ex vivo*, may be implanted directly into the patient without such encapsulation.

Techniques for the encapsulation of living cells are known in the art, and the preparation of the encapsulated cells and their implantation in patients may be routinely accomplished. For example, Baetge et al. (PCT Pub. No. WO 95/05452 and PCT/US94/09299) describe membrane capsules containing genetically engineered cells for the effective delivery of biologically active molecules. The capsules are biocompatible and are easily retrievable. The capsules encapsulate cells transfected with recombinant DNA molecules comprising DNA sequences coding for biologically active molecules operatively linked to promoters that are not subject to down-regulation in vivo upon implantation into a mammalian host.

10

15

20

25

30

The devices provide for the delivery of the molecules from living cells to specific sites within a recipient. In addition, see U.S. Patent Nos. 4,892,538; 5,011,472; and 5,106,627. A system for encapsulating living cells is described in PCT Pub. No. WO 91/10425 (Aebischer et al.). See also, PCT Pub. No. WO 91/10470 (Aebischer et al.); Winn et al., 1991, Exper. Neurol. 113:322-29; Aebischer et al., 1991, Exper. Neurol. 111:269-75; and Tresco et al., 1992, ASAIO 38:17-23.

In vivo and in vitro gene therapy delivery of IL-1ra-R polypeptides is also envisioned. One example of a gene therapy technique is to use the IL-1ra-R gene (either genomic DNA, cDNA, and/or synthetic DNA) encoding an IL-1ra-R polypeptide which may be operably linked to a constitutive or inducible promoter to form a "gene therapy DNA construct." The promoter may be homologous or heterologous to the endogenous IL-1ra-R gene, provided that it is active in the cell or tissue type into which the construct will be inserted. Other components of the gene therapy DNA construct may optionally include DNA molecules designed for site-specific integration (e.g., endogenous sequences useful for homologous recombination), tissue-specific promoters, enhancers or silencers, DNA molecules capable of providing a selective advantage over the parent cell, DNA molecules useful as labels to identify transformed cells, negative selection systems, cell specific binding agents (as, for example, for cell targeting), cell-specific internalization factors, transcription factors enhancing expression from a vector, and factors enabling vector production.

A gene therapy DNA construct can then be introduced into cells (either ex vivo or in vivo) using viral or non-viral vectors. One means for introducing the gene therapy DNA construct is by means of viral vectors as described herein. Certain vectors, such as retroviral vectors, will deliver the DNA construct to the chromosomal DNA of the cells, and the gene can integrate into the chromosomal DNA. Other vectors will function as episomes, and the gene therapy DNA construct will remain in the cytoplasm.

In yet other embodiments, regulatory elements can be included for the controlled expression of the IL-1ra-R gene in the target cell. Such elements are turned on in response to an appropriate effector. In this way, a therapeutic

10

15

20

25

30

polypeptide can be expressed when desired. One conventional control means involves the use of small molecule dimerizers or rapalogs to dimerize chimeric proteins which contain a small molecule-binding domain and a domain capable of initiating a biological process, such as a DNA-binding protein or transcriptional activation protein (*see* PCT Pub. Nos. WO 96/41865, WO 97/31898, and WO 97/31899). The dimerization of the proteins can be used to initiate transcription of the transgene.

An alternative regulation technology uses a method of storing proteins expressed from the gene of interest inside the cell as an aggregate or cluster. The gene of interest is expressed as a fusion protein that includes a conditional aggregation domain that results in the retention of the aggregated protein in the endoplasmic reticulum. The stored proteins are stable and inactive inside the cell. The proteins can be released, however, by administering a drug (e.g., small molecule ligand) that removes the conditional aggregation domain and thereby specifically breaks apart the aggregates or clusters so that the proteins may be secreted from the cell. See Aridor et al., 2000, Science 287:816-17 and Rivera et al., 2000, Science 287:826-30.

Other suitable control means or gene switches include, but are not limited to, the systems described herein. Mifepristone (RU486) is used as a progesterone antagonist. The binding of a modified progesterone receptor ligand-binding domain to the progesterone antagonist activates transcription by forming a dimer of two transcription factors that then pass into the nucleus to bind DNA. The ligand-binding domain is modified to eliminate the ability of the receptor to bind to the natural ligand. The modified steroid hormone receptor system is further described in U.S. Patent No. 5,364,791 and PCT Pub. Nos. WO 96/40911 and WO 97/10337.

Yet another control system uses ecdysone (a fruit fly steroid hormone) which binds to and activates an ecdysone receptor (cytoplasmic receptor). The receptor then translocates to the nucleus to bind a specific DNA response element (promoter from ecdysone-responsive gene). The ecdysone receptor includes a transactivation domain, DNA-binding domain, and ligand-binding domain to

10

15

20

25

30

initiate transcription. The ecdysone system is further described in U.S. Patent No. 5,514,578 and PCT Pub. Nos. WO 97/38117, WO 96/37609, and WO 93/03162.

Another control means uses a positive tetracycline-controllable transactivator. This system involves a mutated tet repressor protein DNA-binding domain (mutated tet R-4 amino acid changes which resulted in a reverse tetracycline-regulated transactivator protein, *i.e.*, it binds to a tet operator in the presence of tetracycline) linked to a polypeptide which activates transcription. Such systems are described in U.S. Patent Nos. 5,464,758, 5,650,298, and 5,654,168.

Additional expression control systems and nucleic acid constructs are described in U.S. Patent Nos. 5,741,679 and 5,834,186, to Innovir Laboratories Inc.

In vivo gene therapy may be accomplished by introducing the gene encoding IL-1ra-R polypeptide into cells via local injection of an IL-1ra-R nucleic acid molecule or by other appropriate viral or non-viral delivery vectors. Hefti, 1994, Neurobiology 25:1418-35. For example, a nucleic acid molecule encoding an IL-1ra-R polypeptide may be contained in an adeno-associated virus (AAV) vector for delivery to the targeted cells (see, e.g., Johnson, PCT Pub. No. WO 95/34670; PCT App. No. PCT/US95/07178). The recombinant AAV genome typically contains AAV inverted terminal repeats flanking a DNA sequence encoding an IL-1ra-R polypeptide operably linked to functional promoter and polyadenylation sequences.

Alternative suitable viral vectors include, but are not limited to, retrovirus, adenovirus, herpes simplex virus, lentivirus, hepatitis virus, parvovirus, papovavirus, poxvirus, alphavirus, coronavirus, rhabdovirus, paramyxovirus, and papilloma virus vectors. U.S. Patent No. 5,672,344 describes an *in vivo* viral-mediated gene transfer system involving a recombinant neurotrophic HSV-1 vector. U.S. Patent No. 5,399,346 provides examples of a process for providing a patient with a therapeutic protein by the delivery of human cells which have been treated *in vitro* to insert a DNA segment encoding a therapeutic protein. Additional methods and materials for the practice of gene therapy techniques are

10

15

20

25

30

described in U.S. Patent Nos. 5,631,236 (involving adenoviral vectors), 5,672,510 (involving retroviral vectors), 5,635,399 (involving retroviral vectors expressing cytokines).

Nonviral delivery methods include, but are not limited to, liposomemediated transfer, naked DNA delivery (direct injection), receptor-mediated transfer (ligand-DNA complex), electroporation, calcium phosphate precipitation, and microparticle bombardment (e.g., gene gun). Gene therapy materials and methods may also include inducible promoters, tissue-specific enhancerpromoters, DNA sequences designed for site-specific integration, DNA sequences capable of providing a selective advantage over the parent cell, labels to identify transformed cells, negative selection systems and expression control systems (safety measures), cell-specific binding agents (for cell targeting), cell-specific internalization factors, and transcription factors to enhance expression by a vector as well as methods of vector manufacture. Such additional methods and materials for the practice of gene therapy techniques are described in U.S. Patent Nos. 4,970,154 (involving electroporation techniques), 5,679,559 (describing a lipoprotein-containing system for gene delivery), 5,676,954 (involving liposome carriers), 5,593,875 (describing methods for calcium phosphate transfection), and 4,945,050 (describing a process wherein biologically active particles are propelled at cells at a speed whereby the particles penetrate the surface of the cells and become incorporated into the interior of the cells), and PCT Pub. No. WO 96/40958 (involving nuclear ligands).

It is also contemplated that IL-1ra-R gene therapy or cell therapy can further include the delivery of one or more additional polypeptide(s) in the same or a different cell(s). Such cells may be separately introduced into the patient, or the cells may be contained in a single implantable device, such as the encapsulating membrane described above, or the cells may be separately modified by means of viral vectors.

A means to increase endogenous IL-1ra-R polypeptide expression in a cell via gene therapy is to insert one or more enhancer elements into the IL-1ra-R polypeptide promoter, where the enhancer elements can serve to increase

10

15

20

25

30

transcriptional activity of the IL-1ra-R gene. The enhancer elements used will be selected based on the tissue in which one desires to activate the gene – enhancer elements known to confer promoter activation in that tissue will be selected. For example, if a gene encoding an IL-1ra-R polypeptide is to be "turned on" in T-cells, the *lck* promoter enhancer element may be used. Here, the functional portion of the transcriptional element to be added may be inserted into a fragment of DNA containing the IL-1ra-R polypeptide promoter (and optionally, inserted into a vector and/or 5' and/or 3' flanking sequences) using standard cloning techniques. This construct, known as a "homologous recombination construct," can then be introduced into the desired cells either *ex vivo* or *in vivo*.

Gene therapy also can be used to decrease IL-1ra-R polypeptide expression by modifying the nucleotide sequence of the endogenous promoter. Such modification is typically accomplished via homologous recombination methods. For example, a DNA molecule containing all or a portion of the promoter of the IL-1ra-R gene selected for inactivation can be engineered to remove and/or replace pieces of the promoter that regulate transcription. For example, the TATA box and/or the binding site of a transcriptional activator of the promoter may be deleted using standard molecular biology techniques; such deletion can inhibit promoter activity thereby repressing the transcription of the corresponding IL-1ra-R gene. The deletion of the TATA box or the transcription activator binding site in the promoter may be accomplished by generating a DNA construct comprising all or the relevant portion of the IL-1ra-R polypeptide promoter (from the same or a related species as the IL-1ra-R gene to be regulated) in which one or more of the TATA box and/or transcriptional activator binding site nucleotides are mutated via substitution, deletion and/or insertion of one or more nucleotides. As a result, the TATA box and/or activator binding site has decreased activity or is rendered completely inactive. This construct, which also will typically contain at least about 500 bases of DNA that correspond to the native (endogenous) 5' and 3' DNA sequences adjacent to the promoter segment that has been modified, may be introduced into the appropriate cells (either ex vivo or in vivo) either directly or via a viral vector as described herein. Typically, the integration of the construct into the genomic DNA of the cells will be via homologous recombination, where the 5' and 3' DNA sequences in the promoter construct can serve to help integrate the modified promoter region via hybridization to the endogenous chromosomal DNA.

5

Therapeutic Uses

IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists thereof can be used to treat, diagnose, ameliorate, or prevent a number of diseases, disorders, or conditions, including those recited herein.

10

15

IL-1ra-R polypeptide agonists and antagonists include those molecules which regulate IL-1ra-R polypeptide activity and either increase or decrease at least one activity of the mature form of the IL-1ra-R polypeptide. Agonists or antagonists may be co-factors, such as a protein, peptide, carbohydrate, lipid, or small molecular weight molecule, which interact with IL-1ra-R polypeptide and thereby regulate its activity. Potential polypeptide agonists or antagonists include antibodies that react with either soluble or membrane-bound forms of IL-1ra-R polypeptides that comprise part or all of the extracellular domains of the said proteins. Molecules that regulate IL-1ra-R polypeptide expression typically include nucleic acids encoding IL-1ra-R polypeptide that can act as anti-sense regulators of expression.

20

25

For example, the IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving immune system dysfunction. Examples of such diseases include, but are not limited to, rheumatoid arthritis, psioriatic arthritis, inflammatory arthritis, osteoarthritis, inflammatory joint disease, autoimmune disease (including autoimmune vasculitis), multiple sclerosis, lupus, diabetes (e.g., insulin diabetes), inflammatory bowel disease, transplant rejection, graft versus host disease, and inflammatory conditions resulting from strain, sprain, cartilage damage, trauma, orthopedic surgery, infection or other disease processes. Other diseases

10

15

20

influenced by the dysfunction of the immune system are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving infection. Examples of such diseases include, but are not limited to, leprosy, viral infections (such as hepatitis or HIV), bacterial infection (such as *clostridium*-associated illnesses, including *clostridium*-associated diarrhea), pulmonary tuberculosis, acute febrile illness, fever, acute phase response of the liver, septicemia, or septic shock. Other diseases involving infection are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving weight disorders. Examples of such diseases include, but are not limited to obesity, anorexia, cachexia (including AIDS-induced cachexia), myopathies (e.g., muscle protein metabolism, such as in sepsis), and hypoglycemia. Other diseases involving weight disorders are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving neuronal dysfunction. Examples of such diseases include, but are not limited to, Alzheimer's disease, Parkinson's disease, neurotoxicity (e.g., as induced by HIV), ALS, brain injury, stress, depression, nociception and other pain (including cancer-related pain), hyperalgesia, epilepsy, learning impairment and memory disorders, sleep disturbance, and peripheral and central neuropathies. Other neurological disorders are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the lung. Examples of such diseases include, but are not limited to, acute or chronic lung injury (including interstitial lung disease), acute respiratory disease syndrome, pulmonary

30

hypertension, emphysema, cystic fibrosis, pulmonary fibrosis, and asthma. Other diseases of the lung are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the skin. Examples of such diseases include, but are not limited to, psoriasis, eczema, and wound healing. Other diseases of the skin are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the kidney. Examples of such diseases include, but are not limited to, acute and chronic glomerulonephritis. Other diseases of the kidney are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the bone. Examples of such diseases include, but are not limited to, osteoporosis, osteopetrosis, osteogenesis imperfecta, Paget's disease, periodontal disease, temporal mandibular joint disease, and hypercalcemia. Other diseases of the bone are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the vascular system. Examples of such diseases include, but are not limited to, hemorrhage or stroke, hemorrhagic shock, ischemia (including cardiac ischemia and cerebral ischemia, e.g., brain injury as a result of trauma, epilepsy, hemorrhage or stroke, each of which may lead to neurodegeneration), atherosclerosis, congestive heart failure, restenosis, reperfusion injury, and angiogenesis. Other diseases of the vascular system are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving tumor cells. Examples of

20

25

30

5

10

10

15

20

25

30

such diseases include, but are not limited to, lymphomas, bone sarcoma, chronic and acute myelogenous leukemia (CML and AML) and other leukemias, multiple myeloma, lung cancer, breast cancer, tumor metastasis, and side effects from radiation therapy. Other diseases involving tumor cells are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the reproductive system. Examples of such diseases include, but are not limited to, infertility, miscarriage, pre-term labor and delivery, and endometriosis. Other diseases involving the reproductive system are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat, diagnose, ameliorate, or prevent diseases, disorders, or conditions involving the eye. Examples of such diseases include, but are not limited to, inflammatory eye disease (as may be associated with, for example, corneal transplant), retinal degeneration, blindness, macular degeneration, glaucoma, uveitis, and retinal neuropathy. Other diseases of the eye are encompassed within the scope of the invention.

The IL-1ra-R nucleic acid molecules, polypeptides, and agonists and antagonists of the invention can also be used to treat diseases such as acute pancreatitis, chronic fatigue syndrome, fibromyalgia, and Kawasaki's disease (MLNS).

IL-1 inhibitors include any protein capable of specifically preventing activation of cellular receptors to IL-1, which may result from any number of mechanisms. Such mechanisms include down-regulating IL-1 production, binding free IL-1, interfering with IL-1 binding to its receptor, interfering with the formation of the IL-1 receptor complex (*i.e.*, association of IL-1 receptor with IL-1 receptor accessory protein), and interfering with the modulation of IL-1 signaling after binding to its receptor. Such interleukin-1 inhibitors include: interleukin-1 receptor antagonists such as IL-1ra-R, as described herein, anti-IL-1 receptor monoclonal antibodies (*e.g.*, European Patent No. 623674), IL-1 binding

proteins such as soluble IL-1 receptors (*e.g.*, U.S. Patent Nos. 5,492,888, 5,488,032, 5,464,937, 5,319,071, and 5,180,812), anti-IL-1 monoclonal antibodies (*e.g.*, PCT Pub. Nos. WO 95/01997, WO 94/02627, WO 90/06371; U.S. Patent No. 4,935,343; and European Patent Nos. 364778, 267611 and 220063), IL-1 receptor accessory proteins and antibodies thereto (*e.g.*, PCT Pub. No. WO 96/23067); inhibitors of interleukin-1E converting enzyme (ICE) or caspase I, which can be used to inhibit IL-1E production and secretion, interleukin-1E protease inhibitors, and other compounds and proteins which block *in vivo* synthesis or extracellular release of IL-1.

10

15

5

Exemplary IL-1 inhibitors are disclosed in US Patent Nos. 5,747,444, 5,359,032, 5,608,035, 5,843,905, 5,359,032, 5,866,576, 5,869,660, 5,869,315, 5,872,095, 5,955,480; PCT Pub. Nos. WO 98/21957, WO 96/09323, WO 91/17184, WO 96/40907, WO 98/32733, WO 98/42325, WO 98/44940, WO 98/47892, WO 98/56377, WO 99/03837, WO 99/06426, WO 99/06042, WO 91/17249, WO 98/32733, WO 98/17661, WO 97/08174, WO 95/34326, WO 99/36426, and WO 99/36415; European Patent Nos. 534978 and 894795; and French Patent Application FR 2762514.

20

25

30

Interleukin-1 receptor antagonist (IL-1ra) is a human protein that acts as a natural inhibitor of interleukin-1. Preferred receptor antagonists (including IL-1ra and variants and derivatives thereof), as well as methods of making and using thereof, are described in U.S. Patent No. 5,075,222; PCT Pub. Nos. WO 91/08285, WO 91/17184, WO 92/16221, WO 93/21946, WO 94/06457, WO 94/21275, WO 94/21235, WO 94/20517, WO 96/22793, WO 97/28828, and WO 99/36541; Austrian Patent No. AU 9173636; French Patent No. FR 2706772; and German Patent No. DE 4219626. Such proteins include glycosylated as well as non-glycosylated IL-1 receptor antagonists.

Specifically, three exemplary forms of IL-1ra and variants thereof are disclosed and described in the 5,075,222 patent. The first of these, called "IL-1i," is characterized as a 22-23 kD molecule on SDS-PAGE with an approximate isoelectric point of 4.8, eluting from a MonoQ FPLC column at around 52 mM NaCl in Tris buffer, pH 7.6. The second, IL-1raE, is characterized as a 22-23 kD

10

15

20

25

30

protein, eluting from a MonoQ column at 48 mM NaCl. Both IL-1ra\Delta and IL-1ra\Delta are glycosylated. The third, IL-1ra\Delta is characterized as a 20 kD protein, eluting from a MonoQ column at 48 mM NaCl, and is non-glycosylated. U.S. Patent No. 5,075,222 also discloses methods for isolating the genes responsible for coding the inhibitors, cloning the gene in suitable vectors and cell types, and expressing the gene to produce the inhibitors.

Those skilled in the art will recognize that many combinations of deletions, insertions, and substitutions (individually or collectively "variant(s)" herein) can be made within the amino acid sequences of IL-1ra-R, provided that the resulting molecule is biologically active (e.g., possesses the ability to affect one or more of the diseases and disorders such as those recited herein.)

As contemplated by the present invention, an IL-1ra-R polypeptide may be administered as an adjunct to other therapy and also with other pharmaceutical compositions suitable for the indication being treated. An IL-1ra-R polypeptide and any of one or more additional therapies or pharmaceutical formulations may be administered separately, sequentially, or simultaneously.

In a specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pre-treatment, post-treatment, or concurrent treatment) with any of one or more TNF inhibitors for the treatment or prevention of the diseases and disorders recited herein.

Such TNF inhibitors include compounds and proteins that block *in vivo* synthesis or extracellular release of TNF. In a specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more of the following TNF inhibitors: TNF binding proteins (soluble TNF receptor type-I and soluble TNF receptor type-II ("sTNFRs"), as defined herein), anti-TNF antibodies, granulocyte colony stimulating factor, thalidomide, BN 50730, tenidap, E 5531, tiapafant PCA 4248, nimesulide, panavir, rolipram, RP 73401, peptide T, MDL 201,449A, (1R,3S)-Cis-1-[9-(2,6-diaminopurinyl)]-3-hydroxy-4-cyclopentene hydrochloride, (1R,3R)-trans-1-(9-(2,6-diamino)purine]-3-acetoxycyclopentane, (1R,3R)-trans-1-[9-adenyl)-3-azidocyclopentane

10

15

20

25

30

hydrochloride and (1R,3R)-trans-1-(6-hydroxy-purin-9-yl)-3-azidocyclo-pentane. TNF binding proteins are disclosed in the art (U.S. Patent No. 5,136,021; European Patent Nos. 308378, 422339, 393438, 398327, 412486, 418014, 433900, 464533, 512528, 526905, 568928, 417563; PCT Pub. Nos. WO 90/13575, WO 91/03553, WO 92/01002, WO 92/13095, WO 92/16221, WO 93/07863, WO 93/21946, WO 93/19777, WO 94/06476, PCT App. No. PCT/US97/12244; English Patent Nos. GB 2218101 and 2246569; and Japanese Patent App. No. JP 127,800/1991).

For example, European Patent Nos. 393438 and 422339 teach the amino acid and nucleic acid sequences of a soluble TNF receptor type I (also known as "sTNFR-I" or "30kDa TNF inhibitor") and a soluble TNF receptor type II (also known as "sTNFR-II" or "40kDa TNF inhibitor"), collectively termed "sTNFRs," as well as modified forms thereof (e.g., fragments, functional derivatives, and variants). European Patent Nos. 393438 and 422339 also disclose methods for isolating the genes responsible for coding the inhibitors, cloning the gene in suitable vectors and cell types, and expressing the gene to produce the inhibitors. Additionally, polyvalent forms (i.e., molecules comprising more than one active moiety) of sTNFR-I and sTNFR-II have also been disclosed. In one embodiment, the polyvalent form may be constructed by chemically coupling at least one TNF inhibitor and another moiety with any clinically acceptable linker, for example polyethylene glycol (PCT Pub. Nos. WO 92/16221 and WO 95/34326), by a peptide linker (Neve et al., 1996, Cytokine, 8:365-70), by chemically coupling to biotin and then binding to avidin (PCT Pub. No. WO 91/03553) and, finally, by combining chimeric antibody molecules (U.S. Patent No. 5,116,964; PCT Pub. Nos. WO 89/09622 and WO 91/16437; and European Patent No. 315062).

Anti-TNF antibodies include MAK 195F Fab antibody (Holler et al., 1993, Ist International Symposium on Cytokines in Bone Marrow Transplantation 147), CDP 571 anti-TNF monoclonal antibody (Rankin et al., 1995, Br. J. Rheumatol., 34:334-42), BAY X 1351 murine anti-tumor necrosis factor monoclonal antibody (Kieft et al., 1995, 7th European Congress of Clinical Microbiology and Infectious Diseases 9); CenTNF cA2 anti-TNF monoclonal

10

15

20

25

30

antibody (Elliott et al., 1994, Lancet, 344:1125-27; Elliott et al., 1994, Lancet, 344:1105-10).

In a specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with secreted or soluble human fas antigen or recombinant versions thereof (PCT Pub. No. WO 96/20206; Mountz *et al.*, 1995, *J. Immunol.*, 155:4829-37; and European Patent No. 510691). PCT Pub. No. WO 96/20206 discloses secreted human fas antigen (native and recombinant, including an Ig fusion protein), methods for isolating the genes responsible for coding the soluble recombinant human fas antigen, methods for cloning the gene in suitable vectors and cell types, and methods for expressing the gene to produce the inhibitors. European Patent No. 510691 teaches nucleic acids coding for human fas antigen, including soluble fas antigen, vectors expressing for said nucleic acids, and transformants transfected with the vector. When administered parenterally, doses of a secreted or soluble fas antigen fusion protein each are generally from about 1 Πg/kg to about 100 Πg/kg.

Current treatment of the diseases and disorders recited herein, including acute and chronic inflammation such as rheumatic diseases, commonly includes the use of first line drugs for control of pain and inflammation; these drugs are classified as non-steroidal, anti-inflammatory drugs (NSAIDs). Secondary treatments include corticosteroids, slow acting antirheumatic drugs (SAARDs), or disease modifying (DM) drugs. Information regarding the following compounds can be found in *The Merck Manual of Diagnosis and Therapy* (16th ed. 1992) and in *Pharmaprojects* (PJB Publications Ltd).

In a specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide and any of one or more NSAIDs for the treatment of the diseases and disorders recited herein, including acute and chronic inflammation such as rheumatic diseases, and graft versus host disease. NSAIDs owe their anti-inflammatory action, at least in part, to the inhibition of prostaglandin synthesis (Goodman and Gilman, *The Pharmacological Basis of Therapeutics* (7th ed. 1985)). NSAIDs can be characterized into at least nine groups: (1) salicylic acid

10

15

20

25

30

derivatives, (2) propionic acid derivatives, (3) acetic acid derivatives, (4) fenamic acid derivatives, (5) carboxylic acid derivatives, (6) butyric acid derivatives, (7) oxicams, (8) pyrazoles, and (9) pyrazolones.

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more salicylic acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. Such salicylic acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof comprise: acetaminosalol, aloxiprin, aspirin, benorylate, bromosaligenin, calcium acetylsalicylate, choline magnesium trisalicylate, magnesium salicylate, choline salicylate, diflusinal, etersalate, fendosal, gentisic acid, glycol salicylate, imidazole salicylate, lysine acetylsalicylate, mesalamine, morpholine salicylate, 1-naphthyl salicylate, olsalazine, parsalmide, phenyl acetylsalicylate, phenyl salicylate, salacetamide, salicylamide O-acetic acid, salsalate, sodium salicylate and sulfasalazine. Structurally related salicylic acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In an additional specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more propionic acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. The propionic acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof comprise: alminoprofen, benoxaprofen, bucloxic acid, carprofen, dexindoprofen, fenoprofen, flunoxaprofen, fluprofen, flurbiprofen, furcloprofen, ibuprofen, ibuprofen aluminum, ibuproxam, indoprofen, isoprofen, ketoprofen, loxoprofen, miroprofen, naproxen, naproxen sodium, oxaprozin, piketoprofen, pimeprofen, pirprofen, pranoprofen, protizinic acid, pyridoxiprofen, suprofen, tiaprofenic acid and tioxaprofen. Structurally related propionic acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In yet another specific embodiment, the present invention is directed to the

10

15

20

25

30

use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more acetic acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. The acetic acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof comprise: acemetacin, alclofenac, amfenac, bufexamac, cinmetacin, clopirac, delmetacin, diclofenac potassium, diclofenac sodium, etodolac, felbinac, fenclofenac, fenclorac, fenclozic acid, fentiazac, furofenac, glucametacin, ibufenac, indomethacin, isofezolac, isoxepac, lonazolac, metiazinic acid, oxametacin, oxpinac, pimetacin, proglumetacin, sulindac, talmetacin, tiaramide, tiopinac, tolmetin, tolmetin sodium, zidometacin and zomepirac. Structurally related acetic acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more fenamic acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. The fenamic acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof comprise: enfenamic acid, etofenamate, flufenamic acid, isonixin, meclofenamic acid, meclofenamate sodium, medofenamic acid, mefenamic acid, niflumic acid, talniflumate, terofenamate, tolfenamic acid and ufenamate. Structurally related fenamic acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In an additional specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more carboxylic acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. The carboxylic acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof which can be used comprise: clidanac, diflunisal, flufenisal, inoridine, ketorolac and tinoridine. Structurally related carboxylic acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

10

15

20

25

30

In yet another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more butyric acid derivatives, prodrug esters, or pharmaceutically acceptable salts thereof. The butyric acid derivatives, prodrug esters, and pharmaceutically acceptable salts thereof comprise: bumadizon, butibufen, fenbufen and xenbucin. Structurally related butyric acid derivatives having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more oxicams, prodrug esters, or pharmaceutically acceptable salts thereof. The oxicams, prodrug esters, and pharmaceutically acceptable salts thereof comprise: droxicam, enolicam, isoxicam, piroxicam, sudoxicam, tenoxicam and 4-hydroxyl-1,2-benzothiazine 1,1-dioxide 4-(N-phenyl)-carboxamide. Structurally related oxicams having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In still another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more pyrazoles, prodrug esters, or pharmaceutically acceptable salts thereof. The pyrazoles, prodrug esters, and pharmaceutically acceptable salts thereof which may be used comprise: difenamizole and epirizole. Structurally related pyrazoles having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In an additional specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment or, concurrent treatment) with any of one or more pyrazolones, prodrug esters, or pharmaceutically acceptable salts thereof. The pyrazolones, prodrug esters, and pharmaceutically acceptable salts thereof which may be used comprise: apazone, azapropazone, benzpiperylon, feprazone, mofebutazone, morazone,

oxyphenbutazone, phenylbutazone, pipebuzone, propylphenazone, ramifenazone, suxibuzone and thiazolinobutazone. Structurally related pyrazalones having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

5

10

15

20

25

30

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more of the following: NSAIDs: H acetamidocaproic acid, S-adenosylmethionine, 3-amino-4-hydroxybutyric acid, amixetrine, anitrazafen, antrafenine, bendazac, bendazac lysinate, benzydamine, ciproquazone, cloximate, bucolome, bufezolac, broperamole, beprozin, dazidamine, deboxamet, detomidine, difenpiramide, difenpyramide, difisalamine, ditazol, emorfazone, fanetizole mesylate, fenflumizole, floctafenine, flumizole, flunixin, fluproquazone, fopirtoline, fosfosal, guaimesal, guaiazolene, isonixirn, lefetamine HCl, leflunomide, lofemizole, lotifazole, lysin clonixinate, meseclazone, nabumetone, nictindole, nimesulide, orgotein, orpanoxin, oxaceprol, oxapadol, paranyline, perisoxal, perisoxal citrate, pifoxime, piproxen, pirazolac, pirfenidone, proquazone, proxazole, thielavin B, tiflamizole, timegadine, tolectin, tolpadol, tryptamid and those designated by company code number such as 480156S, AA861, AD1590, AFP802, AFP860, AI77B, AP504, AU8001, BPPC, BW540C, CHINOIN 127, CN100, EB382, EL508, F1044, FK-506, GV3658, ITF182, KCNTEI6090, KME4, LA2851, MR714, MR897, MY309, ONO3144, PR823, PV102, PV108, R830, RS2131, SCR152, SH440, SIR133, SPAS510, SQ27239, ST281, SY6001, TA60, TAI-901 (4-benzoyl-1-indancarboxylic acid), TVX2706, U60257, UR2301 and WY41770. Structurally related NSAIDs having similar analgesic and anti-inflammatory properties to the NSAIDs are also intended to be encompassed by this group.

In still another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment or concurrent treatment) with any of one or more corticosteroids, prodrug esters, or pharmaceutically acceptable salts thereof for the treatment of the diseases and disorders recited herein, including acute and chronic inflammation such as

10

15

20

25

30

rheumatic diseases, graft versus host disease, and multiple sclerosis. Corticosteroids, prodrug esters, and pharmaceutically acceptable salts thereof include hydrocortisone and compounds which are derived from hydrocortisone, amcinonide, algestone, such as 21-acetoxypregnenolone, alclomerasone, budesonide. betamethasone valerate, betamethasone, beclomethasone, chloroprednisone, clobetasol, clobetasol propionate, clobetasone, clobetasone cloprednol, corticosterone, cortisone, cortivazol, clocortolone, butyrate, diflorasone, dexamethasone, desoximerasone, desonide, deflazacon, diflucortolone, difluprednate, enoxolone, fluazacort, flucloronide, flumethasone, flucinolone acetonide, flunisolide, fluocinonide. flumethasone pivalate, fluorocinolone acetonide, fluocortin butyl, fluocortolone, fluocortolone hexanoate, diflucortolone valerate, fluorometholone, fluperolone acetate, fluprednidene acetate, fluprednisolone, flurandenolide, formocortal, halcinonide, halometasone, halopredone acetate, hydrocortamate, hydrocortisone, hydrocortisone acetate, hydrocortisone butyrate, hydrocortisone phosphate, hydrocortisone 21-sodium succinate, hydrocortisone tebutate, mazipredone, medrysone, meprednisone, paramethasone, prednicarbate, furoate, mometasone methylprednisolone, prednisolone sodium 21-diedryaminoacetate, prednisolone prednisolone, phosphate, prednisolone sodium succinate, prednisolone sodium 21-msulfobenzoate, prednisolone sodium 21-stearoglycolate, prednisolone tebutate, prednival, prednylidene, prednisone, 21-trimethylacetate, prednisolone prednylidene 21-diethylaminoacetate, tixocortol, triamcinolone, triamcinolone acetonide, triamcinolone benetonide and triamcinolone hexacetonide. Structurally related corticosteroids having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more slow-acting antirheumatic drugs (SAARDs) or disease modifying antirheumatic drugs (DMARDS), prodrug esters, or pharmaceutically acceptable salts thereof for the treatment of the diseases and disorders recited herein, including acute and chronic inflammation such as

rheumatic diseases, graft versus host disease, and multiple sclerosis. SAARDs or DMARDS, prodrug esters, and pharmaceutically acceptable salts thereof comprise: allocupreide sodium, auranofin, aurothioglucose, aurothioglycanide, azathioprine, brequinar sodium, bucillamine, calcium 3-aurothio-2-propanol-1-sulfonate, chlorambucil, chloroquine, clobuzarit, cuproxoline, cyclophosphamide, cyclosporin, dapsone, 15-deoxyspergualin, diacerein, glucosamine, gold salts (e.g., cycloquine gold salt, gold sodium thiomalate, gold sodium thiosulfate), hydroxychloroquine, hydroxychloroquine sulfate, hydroxyurea, kebuzone, levamisole, lobenzarit, melittin, 6-mercaptopurine, methotrexate, mizoribine, mycophenolate mofetil, myoral, nitrogen mustard, D-penicillamine, pyridinol imidazoles such as SKNF86002 and SB203580, rapamycin, thiols, thymopoietin and vincristine. Structurally related SAARDs or DMARDs having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

15

20

25

30

10

5

In another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more COX2 inhibitors, prodrug esters, or pharmaceutically acceptable salts thereof for the treatment of the diseases and disorders recited herein, including acute and chronic inflammation. Examples of COX2 inhibitors, prodrug esters, or pharmaceutically acceptable salts thereof include, for example, celecoxib. Structurally related COX2 inhibitors having similar analgesic and anti-inflammatory properties are also intended to be encompassed by this group.

In still another specific embodiment, the present invention is directed to the use of an IL-1ra-R polypeptide in combination (pretreatment, post-treatment, or concurrent treatment) with any of one or more antimicrobials, prodrug esters, or pharmaceutically acceptable salts thereof for the treatment of the diseases and chronic inflammation. acute and disorders recited herein, including Antimicrobials include, for example, the broad classes of penicillins, cephalosporins and other beta-lactams, aminoglycosides, azoles, quinolones, lincosamides and sulfonamides, tetracyclines, macrolides, rifamycins,

10

15

20

25

30

polymyxins. The penicillins include, but are not limited to, penicillin G, penicillin V, methicillin, nafcillin, oxacillin, cloxacillin, dicloxacillin, floxacillin, ampicillin, ampicillin/sulbactam, amoxicillin, amoxicillin/clavulanate, hetacillin, cyclacillin, ticarcillin, indanyl, bacampicillin, carbenicillin, carbenicillin ticarcillin/clavulanate, azlocillin, mezlocillin, peperacillin, and mecillinam. The cephalosporins and other beta-lactams include, but are not limited to, cephalothin, cephapirin, cephalexin, cephradine, cefazolin, cefadroxil, cefaclor, cefamandole, cefotetan, cefoxitin, ceruroxime, cefonicid, ceforadine, cefixime, cefotaxime, moxalactam, ceftizoxime, cetriaxone, cephoperazone, ceftazidime, imipenem and aztreonam. The aminoglycosides include, but are not limited to, streptomycin, gentamicin, tobramycin, amikacin, netilmicin, kanamycin and neomycin. The azoles include, but are not limited to, fluconazole. The quinolones include, but are not limited to, nalidixic acid, norfloxacin, enoxacin, ciprofloxacin, ofloxacin, sparfloxacin and temafloxacin. The macrolides include, but are not limited to, erythomycin, spiramycin and azithromycin. The rifamycins include, but are not limited to, rifampin. The tetracyclines include, but are not limited to, spicycline, chlortetracycline, clomocycline, demeclocycline, deoxycycline, guamecycline, lymecycline, meclocycline, methacycline, minocycline, oxytetracycline, penimepicycline, pipacycline, rolitetracycline, sancycline, senociclin and tetracycline. The sulfonamides include, but are not limited to, sulfanilamide, sulfamethoxazole, sulfacetamide, sulfadiazine, sulfisoxazole and co-trimoxazole (trimethoprim/sulfamethoxazole). The lincosamides include, but are not limited to, clindamycin and lincomycin. The polymyxins (polypeptides) include, but are not limited to, polymyxin B and colistin.

Agonists or antagonists of IL-1RA-R polypeptide function may be used (simultaneously or sequentially) in combination with one or more cytokines, growth factors, anti-inflammatories, and/or chemotherapeutic agents as is appropriate for the condition being treated.

Other diseases caused by or mediated by undesirable levels of one or more of IL-1, IL-1ra, or IL-1ra-R polypeptide are encompassed within the scope of the

invention. Undesirable levels include excessive levels of IL-1, IL-1ra, or IL-1ra-R polypeptide and sub-normal levels of IL-1, IL-1ra, or IL-1ra-R polypeptide.

Uses of IL-1ra-R Nucleic Acids and Polypeptides

Nucleic acid molecules of the invention (including those that do not themselves encode biologically active polypeptides) may be used to map the locations of the IL-1ra-R gene and related genes on chromosomes. Mapping may be done by techniques known in the art, such as PCR amplification and *in situ*

hybridization.

5

10

15

20

IL-1ra-R nucleic acid molecules (including those that do not themselves encode biologically active polypeptides), may be useful as hybridization probes in diagnostic assays to test, either qualitatively or quantitatively, for the presence of an IL-1ra-R nucleic acid molecule in mammalian tissue or bodily fluid samples.

Other methods may also be employed where it is desirable to inhibit the activity of one or more IL-1ra-R polypeptides. Such inhibition may be effected by nucleic acid molecules that are complementary to and hybridize to expression control sequences (triple helix formation) or to IL-1ra-R mRNA. For example, antisense DNA or RNA molecules, which have a sequence that is complementary to at least a portion of an IL-1ra-R gene can be introduced into the cell. Antisense probes may be designed by available techniques using the sequence of the IL-1ra-R gene disclosed herein. Typically, each such antisense molecule will be complementary to the start site (5' end) of each selected IL-1ra-R gene. When the antisense molecule then hybridizes to the corresponding IL-1ra-R mRNA, translation of this mRNA is prevented or reduced. Anti-sense inhibitors provide information relating to the decrease or absence of an IL-1ra-R polypeptide in a cell or organism.

Alternatively, gene therapy may be employed to create a dominant-negative inhibitor of one or more IL-1ra-R polypeptides. In this situation, the DNA encoding a mutant polypeptide of each selected IL-1ra-R polypeptide can be prepared and introduced into the cells of a patient using either viral or non-viral methods as described herein. Each such mutant is typically designed to compete

30

10

15

20

25

30

with endogenous polypeptide in its biological role.

In addition, an IL-1ra-R polypeptide, whether biologically active or not, may be used as an immunogen, that is, the polypeptide contains at least one epitope to which antibodies may be raised. Selective binding agents that bind to an IL-1ra-R polypeptide (as described herein) may be used for *in vivo* and *in vitro* diagnostic purposes, including, but not limited to, use in labeled form to detect the presence of IL-1ra-R polypeptide in a body fluid or cell sample. The antibodies may also be used to prevent, treat, or diagnose a number of diseases and disorders, including those recited herein. The antibodies may bind to an IL-1ra-R polypeptide so as to diminish or block at least one activity characteristic of an IL-1ra-R polypeptide, or may bind to a polypeptide to increase at least one activity characteristic of an IL-1ra-R polypeptide (including by increasing the pharmacokinetics of the IL-1ra-R polypeptide).

The IL-1ra-R polypeptides of the present invention can be used to clone IL-1ra-R polypeptide receptors, using an expression cloning strategy. Radiolabeled (125 Iodine) IL-1ra-R polypeptide or affinity/activity-tagged IL-1ra-R polypeptide (such as an Fc fusion or an alkaline phosphatase fusion) can be used in binding assays to identify a cell type or cell line or tissue that expresses IL-1ra-R polypeptide receptors. RNA isolated from such cells or tissues can be converted to cDNA, cloned into a mammalian expression vector, and transfected into mammalian cells (such as COS or 293 cells) to create an expression library. A radiolabeled or tagged IL-1ra-R polypeptide can then be used as an affinity ligand to identify and isolate from this library the subset of cells that express the IL-1ra-R polypeptide receptors on their surface. DNA can then be isolated from these cells and transfected into mammalian cells to create a secondary expression library in which the fraction of cells expressing IL-1ra-R polypeptide receptors is many-fold higher than in the original library. This enrichment process can be repeated iteratively until a single recombinant clone containing an IL-1ra-R polypeptide receptor is isolated. Isolation of the IL-1ra-R polypeptide receptors is useful for identifying or developing novel agonists and antagonists of the IL-1ra-R polypeptide signaling pathway. Such agonists and antagonists include soluble

10

15

20

25

30

IL-1ra-R polypeptide receptors, anti-IL-1ra-R polypeptide receptor antibodies, small molecules, or antisense oligonucleotides, and they may be used for treating, preventing, or diagnosing one or more of the diseases or disorders described herein.

A deposit of cDNA encoding human IL-1ra-R polypeptide, having Accession No. PTA-1423, was made with the American Type Culture Collection (ATCC), 10801 University Boulevard, Manassas, VA 20110-2209 on February 29, 2000.

The following examples are intended for illustration purposes only, and should not be construed as limiting the scope of the invention in any way.

Example 1: Cloning of the Human IL-1ra-R Polypeptide Gene

Generally, materials and methods as described in Sambrook et al. supra were used to clone and analyze the gene encoding human IL-1ra-R polypeptide.

To isolate cDNA sequences encoding human IL-1ra-R polypeptide, a search of a proprietary database (Amgen, Thousand Oaks, CA) was performed using clone shos1-00003-d1 as a query sequence. A 423 bp sequence identified in this manner was found to share 41% identity to the query sequence at the amino acid level. The region of highest homology within this sequence was used to design gene specific oligonucleotides for the identification of cDNA sources and the generation of cDNA clones, using various PCR strategies.

A number of cDNA libraries were analyzed in amplification reactions containing 10ng of cDNA library template DNA, 10 pmol each of the amplimers 2349-98 (5'-C-A-C-A-C-G-C-T-T-C-A-C-C-T-T-C-T-T-C-C-A-G-3'; SEQ ID NO: 20) and 2349-99 (5'-T-A-A-A-A-C-T-T-G-G-T-A-C-G-G-C-T-G-A-G-G-G-3'; SEQ ID NO: 21), and Ready-To-Go PCR beads (Amersham-Pharmacia, Piscataway, NJ) (Pharmacia, Piscataway, NJ), in a total reaction volume of 25 Π . Reactions were performed at 95°C for 5 minutes for one cycle; 95°C for 15 seconds, 63°C for 15 seconds, and 72°C for 1 minute for 30 cycles; and 72°C for 10 minutes for one cycle. A PCR product of the expected size (153 bp) was identified in a number of cDNA libraries, including human fetal scalp (oligo-dT

primed) and human placenta (oligo-dT primed and random primed).

The fetal scalp and placenta cDNA libraries were prepared as follows. Total RNA was extracted from human fetal scalp and from human placenta using standard RNA extraction procedures and poly-A⁺ RNA was selected from this total RNA using standard procedures. Random primed or oligo-dT primed cDNA was synthesized from this poly-A⁺ RNA using the Superscript Plasmid System for cDNA Synthesis and Plasmid Cloning kit (Gibco-BRL), according to the manufacturer's suggested protocols, or other suitable procedure. The resulting cDNA was digested with appropriate restriction enzymes and was then ligated into pSPORT-1, or other suitable cloning vector. Ligation products were transformed into *E. coli* using standard techniques, and bacterial transformants were selected on culture plates containing either ampicillin, tetracycline, kanamycin, or chloramphenicol. The cDNA library consisted of all, or a subset, of these transformants.

15

20

25

30

10

5

Both 5'RACE and 3'RACE reactions were performed in order to generate the full-length cDNA sequence for IL-1ra-R polypeptide. To isolate cDNA sequences corresponding to the 5' end of the cDNA sequence for IL-1ra-R polypeptide, 5'RACE was performed using 10 ng of an oligo-dT-primed human fetal scalp cDNA library in pSPORT1 and the primers 1572-36 (5'-G-T-G-T-G-G-A-A-T-T-G-T-G-A-G-C-G-G-A-T-A-A-C-3'; SEQ ID NO: 22) and 2349-99. Reactions were performed at 94°C for 1 minute for one cycle; 94°C for 5 seconds and 72°C for 5 minutes for 5 cycles; 94°C for 5 seconds, 70°C for 10 seconds, and 72°C for 3 minutes for 5 cycles; 94°C for 5 seconds, 68°C for 10 seconds, and 72°C for 3 minutes for 25 cycles; and 72°C for 7 minutes for one cycle. Nested PCR was performed using a portion of the 5'RACE amplification product and the primers 2328-91 (5'-C-T-A-T-G-A-C-C-A-T-G-A-T-T-A-C-G-C-C-A-A-G-C-3'; SEQ ID NO: 23) and 2351-47 (5'-G-C-T-G-T-A-C-T-G-G-C-T-G-C-T-G-G-G-G-C-3'; SEQ ID NO: 24). Nested PCR was performed at 94°C for 1 minute for one cycle; 94°C for 5 seconds and 72°C for 5 minutes for 5 cycles; 94°C for 5 seconds, 70°C for 10 seconds, and 72°C for 3 minutes for 5 cycles; 94°C for 5

10

15

20

25

30

seconds, 68°C for 10 seconds, and 72°C for 3 minutes for 25 cycles; and 72°C for 7 minutes for one cycle.

To isolate cDNA sequences corresponding to the 3' end of the cDNA sequence for IL-1ra-R polypeptide, 3'RACE was performed using 10 ng of an oligo-dT-primed human fetal scalp cDNA library in pSPORT1 and the primers 2351-48 (5'-C-C-T-T-C-A-G-G-C-T-T-G-A-G-G-C-T-G-3'; SEQ ID NO: 25) and 2329-93 (5'-C-G-G-G-C-C-T-C-T-T-C-G-C-T-A-T-T-A-C-G-C-3'; SEQ ID NO: 26). Reactions were performed at 94°C for 1 minute for one cycle; 94°C for 5 seconds and 72°C for 5 minutes for 5 cycles; 94°C for 5 seconds, 70°C for 10 seconds, and 72°C for 3 minutes for 5 cycles; 94°C for 5 seconds, 68°C for 10 seconds, and 72°C for 3 minutes for 25 cycles; and 72°C for 7 minutes for one cycle. Nested PCR was performed using a portion of the 3'RACE amplification G-G-C-3'; SEQ ID NO: 27) and 2329-94 (5'-T-G-G-C-G-A-A-A-G-G-G-G-G-G-A-T-G-T-G-C-T-G-3'; SEQ ID NO: 28). Nested PCR was performed at 95°C for 1 minute for one cycle; 95°C for 30 seconds and 68°C for 1 minute for 30 cycles; and 68°C for 7 minutes for one cycle. The full-length cDNA sequence for IL-1ra-R polypeptide was assembled from the resulting collection of 5'RACE and 3'RACE clones.

The 5' portion of the cDNA sequence for IL-1ra-R polypeptide was confirmed by independently isolating cDNA sequences from a human placenta Marathon™ cDNA library (Clontech). To isolate cDNA sequences corresponding to the 5' end of the cDNA sequence for IL-1ra-R polypeptide, 5'RACE was performed using 1 ng of human placenta cDNA and the primers AP-1 (5'-C-C-A-T-C-T-A-A-T-A-C-G-A-C-T-C-A-C-T-A-T-A-G-G-G-C-3'; SEQ ID NO: 29; Clontech) and 2353-87 (5'-C-C-T-T-G-G-T-G-A-G-C-T-G-T-A-C-T-G-G-C-T-G-3'; SEQ ID NO: 30). Reactions were performed at 94°C for 2 minutes for one cycle; 94°C for 5 seconds and 72°C for 1.5 minutes for 5 cycles; 94°C for 5 seconds and 68°C for 1.5 minutes for 25 cycles; and 86°C for 7 minutes for one cycle. Nested PCR was performed using a portion of the 5'RACE amplification product and the primers

10

15

20

25

30

AP-1 and 2349-52 (5'-C-C-G-G-C-C-A-C-A-C-A-G-G-A-A-C-C-A-3'; SEQ ID NO: 31). Nested PCR was performed at 94°C for 2 minutes for one cycle; 94°C for 10 seconds and 68°C for 1.5 minutes for 30 cycles; and 68°C for 7 minutes for one cycle.

Sequence analysis of the predicted cDNA sequence for human IL-1ra-R polypeptide indicated that the gene comprises a 456 bp open reading frame encoding a protein of 152 amino acids (Figures 1A-1B). A sequence variant of the human IL-1ra-R polypeptide was also identified. Sequence analysis of this variant indicated that the gene for this variant also comprises a 456 bp open reading frame encoding a protein of 152 amino acids (Figures 2A-2B). The nucleotide sequence of this variant differs from that of the human IL-1ra-R gene at two nucleotide positions.

Figures 4A-4B illustrate the amino acid sequence alignment of human IL-1Δ (IL-1_alpha; SEQ ID NO: 7), human IL-1E (IL-1_beta; SEQ ID NO: 8), human IL-1 receptor antagonist (IL-1RA; SEQ ID NO: 9), human IL-1Γ (IL-1_delta; SEQ ID NO: 10), human IL-1ra-R polypeptide (IL-1ra-R; SEQ ID NO: 2), human Tango-77 (Tango-77; SEQ ID NO: 11), human Zilla4 (Zilla4; SEQ ID NO: 12), human IL-1] (IL-1_zeta; SEQ ID NO: 13), human IL-1 receptor antagonist E (IL-1RA_beta; SEQ ID NO: 14), human SPOIL II (Spoil_II; SEQ ID NO: 15), human IL-1H (IL-1_epsilon; SEQ ID NO: 16), and human IL-1K (IL-1_eta; SEQ ID NO: 17). Figure 5 schematically illustrates the phylogenetic relationship of the IL-1ra gene family.

Example 2: Cloning of the Human IL-1ra-R Polypeptide Gene Splice Variant

Generally, materials and methods as described in Sambrook *et al. supra* were used to clone and analyze the gene encoding a human IL-1ra-R polypeptide splice variant.

Full-length cDNA sequences encoding a splice variant of human IL-1ra-R polypeptide, were isolated from a human placenta cDNA library, which was prepared as follows. Total RNA was extracted from human placenta using standard RNA extraction procedures and poly-A⁺ RNA was selected from this

10

15

20

25

30

total RNA using standard procedures. Oligo-dT primed cDNA was synthesized from this poly-A⁺ RNA using the Superscript Plasmid System for cDNA Synthesis and Plasmid Cloning kit (Gibco-BRL), according to the manufacturer's suggested protocols. The resulting cDNA was digested with Not I and then fractionated by electrophoresis on a 0.8% agarose gel. Fragments of 0.8 to 1.6 kb were isolated, purified, and then ligated into pSPORT-1. Ligation products were transformed into *E. coli* using standard techniques, and bacterial transformants were selected on culture plates containing ampicillin. The resulting transformants were pooled to generate the cDNA library and plasmid DNA was prepared from 12 pools (each containing approximately 80,000 colonies) of this cDNA library.

Each of the cDNA library pools was analyzed in amplification reactions containing 10ng of template DNA, the amplimers 2349-98 and 2349-99 at a concentration of 0.4 TM each, and Ready-To-Go PCR beads, in a total reaction volume of 25 II. Reactions were performed at 94°C for 5 minutes for one cycle; 94°C for 30 seconds, 65°C for 30 seconds, and 72°C for 1 minute for 30 cycles; and 72°C for 10 minutes for one cycle. The cDNA library pools were also analyzed in amplification reactions containing 10ng of template DNA, the amplimers 2349-51 (5'-A-A-G-A-G-C-C-A-C-A-C-G-C-T-T-C-A-C-C-T-T-C-T-3'; SEQ ID NO: 32) and 2349-52 at a concentration of 0.4 IM each, and Ready-To-Go PCR beads, in a total reaction volume of 25 II. Reactions were performed at 94°C for 5 minutes for one cycle; 94°C for 30 seconds, 65°C for 30 seconds, and 72°C for 1 minute for 30 cycles; and 72°C for 10 minutes for one cycle. The PCR products obtained in these reactions were then analyzed on agarose gels. Plasmid DNA from three pools yielded PCR products having the expected size, and 3 x 10⁵ clones from each of these positive pools were screened by PCR using the amplimers 2349-98 and 2349-99. One of the PCR products (zhvt-000329) generated by PCR was labeled with 32P-dCTP and used as a probe to re-screen the positive cDNA library pools. Bacteria colonies were plated (at approximately 50,000 per 150 mm plate) and then lifted onto nitrocellulose filters. Filters were prehybridized in ExpressHyb hybridization solution (Clontech) for 30 minutes at 68°C, and then hybridized in the same solution with the addition of

10

15

20

25

30

labeled probe at 68°C overnight. Following hybridization, the filters were washed twice for 10 minutes at room temperature in 2X SSC and 0.05% SDS, and then twice for 30 minutes at 68 °C in 0.1X SSC and 0.1% SDS. Filters were then exposed to autoradiography with intensifying screens at -80°C for 2 hours. A positive clone (RDS#199918503) containing an insert of approximately 1.2 kb was identified from one of the cDNA library pools. A second positive clone (RDS#199918501) containing insert size about 0.8 kb was identified from a separate cDNA library pool. Plasmid DNA was prepared from each of these clones and the sequence was analyzed.

Sequence analysis of the predicted cDNA sequence for one of these clones indicated that the cDNA encoded a variant of human IL-1ra-R polypeptide (see Example 1). The gene encoding this variant comprises a 513 bp open reading frame encoding a protein of 171 amino acids (Figure 3).

Sequence analysis of the human genomic sequence containing the IL-1ra-R gene (Genbank Accession No. AC016724, contig. fragment 76649-96342) indicated that the first exon of the IL-1ra-R gene (see Example 1) lies 4.1 kb downstream of the last exon of IL-1 Omega (Genbank Accession No. Z300050). The close proximity of these two genes in the genome suggests that the present polypeptide might arise from splicing (or a fusion) of the first two exons of IL-1 Omega onto the second exon of the IL-1ra-R gene, or a variant thereof (see Example 1). This variant of the IL-1ra-R polypeptide may therefore be an IL-1ra-R splice variant, fusion protein, or the like. Juxtaposition of the second exon in IL-1 Omega and the second exon of the IL-1ra-R gene (or a variant thereof) results in a sequence that encodes a protein with an N-terminus that appears to function as a signal peptide. Figure 6 schematically illustrates the relationship between human IL-1ra-R polypeptide (Mature CS329), the sequence variant of human IL-1ra-R polypeptide (Mature CS329 Variant protein), and the splice variant of human IL-1ra-R polypeptide (Omega 329 protein).

10

15

20

Generally, materials and methods as described in Sambrook et al. supra were used to clone and analyze the gene encoding murine IL-1ra-R polypeptide.

To isolate cDNA sequences encoding murine IL-1ra-R polypeptide, PCR was performed using a day 7 mouse embryo cDNA library template and the amplimers 2557-95 (5'-A-A-G-C-C-T-T-T-T-T-C-T-T-C-T-T-G-C-C-T-C-A-G-T-G-3'; SEQ ID NO: 33) and 2557-96 (5'-T-G-C-C-A-T-T-T-A-A-T-G-T-A-A-C-A-C-G-G-T-C-A-C-A-G-3'; SEQ ID NO: 34) and standard techniques.

Sequence analysis of the predicted cDNA sequence for murine IL-1ra-R polypeptide indicated that the gene comprises a 456 bp open reading frame encoding a protein of 152 amino acids (Figure 7). Figure 8 illustrates the amino acid sequence alignment of human IL-1ra-R polypeptide (huIL-1ra-R; SEQ ID NO: 2) and murine IL-1ra-R polypeptide (muIL-1ra-R; SEQ ID NO: 36). Figures 9A-9I illustrate the genomic nucleotide sequence for the murine IL-1ra-R gene (SEQ ID NO: 37). The locations of the coding portions of exons 1-4 are indicated (underline).

Example 4: IL-1ra-R mRNA Expression

Expression of IL-1ra-R mRNA was examined by PCR in amplification reactions containing 10ng of cDNA library template DNA, 10 pmol of the amplimers 2349-98 and 2349-99, and Ready-To-Go PCR beads, in a total reaction volume of 25 II. Reactions were performed at 95°C for 5 minutes for one cycle; 95°C for 15 seconds, 63°C for 15 seconds, and 72°C for 1 minute for 30 cycles; and 72°C for 10 minutes for one cycle. A PCR product of the expected size (153 bp) was identified in a number of cDNA libraries, including human fetal scalp, human fetal eye, human gall bladder, and human placenta. IL-1ra-R mRNA expression was also detected in fetal eye and fetal spleen by RT-PCR using the amplimers 2349-51 and 2349-52 and the Titan system (Boehringer). These reactions were performed at 55°C for 30 minutes for one cycle; 94°C for 15 seconds, 64°C for 15 seconds, and 68°C for 50 seconds (with an increase of 2 seconds per cycle) for 30 cycles; and 68°C for 7 minutes for one cycle.

30

10

15

20

25

30

Expression of IL-1ra-R mRNA was examined by Northern blot analysis using human and mouse RNA blots (Clontech). Blots were first prehybridized for 30 minutes at 65°C in Pre-Hyb Solution (Amersham) and then probed overnight at 65°C in the same solution containing 25 ng of a 32P-labeled probe corresponding to nucleotides 1-474 of the cDNA sequence encoding IL-1ra-R polypeptide. The probe was labeled using a Redi Prime II kit (Pharmacia). Following hybridization, blots were washed twice for 1 hour in 6X SSC and 0.1% SDS, twice for 1 hour in 2X SSC and 0.1% SDS, twice for 30 minutes in 0.2X SSC and 0.1% SDS, and then were rinsed in 2X SSC. Following overnight exposure of the Northern blots in a phosphoimager, a transcript of approximately 1.6 kb was detected in mouse skeletal muscle (Figure 10A) and a transcript of approximately 2.9 kb was detected in human pancreas and peripheral blood leukocytes (Figure 10B and 10C). In an analysis of dot blots (Clontech) using the same probe, IL-1ra-R mRNA was detected in murine skeletal muscle, submaxillar gland, and epididymis.

The expression of IL-1ra-R mRNA is localized by *in situ* hybridization. A panel of normal embryonic and adult mouse tissues is fixed in 4% paraformaldehyde, embedded in paraffin, and sectioned at 5 Tm. Sectioned tissues are permeabilized in 0.2 M HCl, digested with Proteinase K, and acetylated with triethanolamine and acetic anhydride. Sections are prehybridized for 1 hour at 60°C in hybridization solution (300 mM NaCl, 20 mM Tris-HCl, pH 8.0, 5 mM EDTA, 1X Denhardt's solution, 0.2% SDS, 10 mM DTT, 0.25 mg/ml tRNA, 25 Tg/ml polyA, 25 Tg/ml polyC and 50% formamide) and then hybridized overnight at 60°C in the same solution containing 10% dextran and 2 x 10⁴ cpm/Tl of a ³³P-labeled antisense riboprobe complementary to the human IL-1ra-R gene. The riboprobe is obtained by *in vitro* transcription of a clone containing human IL-1ra-R cDNA sequences using standard techniques.

Following hybridization, sections are rinsed in hybridization solution, treated with RNaseA to digest unhybridized probe, and then washed in 0.1X SSC at 55°C for 30 minutes. Sections are then immersed in NTB-2 emulsion (Kodak,

10

15

20

Rochester, NY), exposed for 3 weeks at 4°C, developed, and counterstained with hematoxylin and eosin. Tissue morphology and hybridization signal are simultaneously analyzed by darkfield and standard illumination for brain (one sagittal and two coronal sections), gastrointestinal tract (esophagus, stomach, duodenum, jejunum, ileum, proximal colon, and distal colon), pituitary, liver, lung, heart, spleen, thymus, lymph nodes, kidney, adrenal, bladder, pancreas, salivary gland, male and female reproductive organs (ovary, oviduct, and uterus in the female; and testis, epididymus, prostate, seminal vesicle, and vas deferens in the male), BAT and WAT (subcutaneous, peri-renal), bone (femur), skin, breast, and skeletal muscle.

Example 5: Production of IL-1ra-R Polypeptides

A. Expression of IL-1ra-R Polypeptides in Bacteria

PCR is used to amplify template DNA sequences encoding an IL-1ra-R polypeptide using primers corresponding to the 5' and 3' ends of the sequence. The amplified DNA products may be modified to contain restriction enzyme sites to allow for insertion into expression vectors. PCR products are gel purified and inserted into expression vectors using standard recombinant DNA methodology. An exemplary vector, such as pAMG21 (ATCC no. 98113) containing the lux promoter and a gene encoding kanamycin resistance is digested with Bam HI and Nde I for directional cloning of inserted DNA. The ligated mixture is transformed into an *E. coli* host strain by electroporation and transformants are selected for kanamycin resistance. Plasmid DNA from selected colonies is isolated and subjected to DNA sequencing to confirm the presence of the insert.

Transformed host cells are incubated in 2xYT medium containing 30 Π_g/mL kanamycin at 30°C prior to induction. Gene expression is induced by the addition of N-(3-oxohexanoyl)-dl-homoserine lactone to a final concentration of 30 ng/mL followed by incubation at either 30°C or 37°C for six hours. The expression of IL-1ra-R polypeptide is evaluated by centrifugation of the culture, resuspension and lysis of the bacterial pellets, and analysis of host cell proteins by SDS-polyacrylamide gel electrophoresis.

30

10

15

20

Inclusion bodies containing IL-1ra-R polypeptide are purified as follows. Bacterial cells are pelleted by centrifugation and resuspended in water. The cell suspension is lysed by sonication and pelleted by centrifugation at 195,000 xg for 5 to 10 minutes. The supernatant is discarded, and the pellet is washed and transferred to a homogenizer. The pellet is homogenized in 5 mL of a Percoll solution (75% liquid Percoll and 0.15 M NaCl) until uniformly suspended and then diluted and centrifuged at 21,600 xg for 30 minutes. Gradient fractions containing the inclusion bodies are recovered and pooled. The isolated inclusion bodies are analyzed by SDS-PAGE.

A single band on an SDS polyacrylamide gel corresponding to *E. coli*-produced IL-1ra-R polypeptide is excised from the gel, and the N-terminal amino acid sequence is determined essentially as described by Matsudaira *et al.*, 1987, *J. Biol. Chem.* 262:10-35.

B. Expression of IL-1ra-R Polypeptide in Mammalian Cells

PCR is used to amplify template DNA sequences encoding an IL-1ra-R polypeptide using primers corresponding to the 5' and 3' ends of the sequence. The amplified DNA products may be modified to contain restriction enzyme sites to allow for insertion into expression vectors. PCR products are gel purified and inserted into expression vectors using standard recombinant DNA methodology. An exemplary expression vector, pCEP4 (Invitrogen, Carlsbad, CA), that contains an Epstein-Barr virus origin of replication, may be used for the expression of IL-1ra-R polypeptides in 293-EBNA-1 cells. Amplified and gel purified PCR products are ligated into pCEP4 vector and introduced into 293-EBNA cells by lipofection. The transfected cells are selected in 100 Fg/mL hygromycin and the resulting drug-resistant cultures are grown to confluence. The cells are then cultured in serum-free media for 72 hours. The conditioned media is removed and IL-1ra-R polypeptide expression is analyzed by SDS-PAGE.

IL-1ra-R polypeptide expression may be detected by silver staining. Alternatively, IL-1ra-R polypeptide is produced as a fusion protein with an

30

10

15

20

25

epitope tag, such as an IgG constant domain or a FLAG epitope, which may be detected by Western blot analysis using antibodies to the peptide tag.

IL-1ra-R polypeptides may be excised from an SDS-polyacrylamide gel, or IL-1ra-R fusion proteins are purified by affinity chromatography to the epitope tag, and subjected to N-terminal amino acid sequence analysis as described herein.

C. Expression and Purification of IL-1ra-R Polypeptide in Mammalian Cells

IL-1ra-R polypeptide expression constructs are introduced into 293 EBNA or CHO cells using either a lipofection or calcium phosphate protocol.

To conduct functional studies on the IL-1ra-R polypeptides that are produced, large quantities of conditioned media are generated from a pool of hygromycin selected 293 EBNA clones. The cells are cultured in 500 cm Nunc Triple Flasks to 80% confluence before switching to serum free media a week prior to harvesting the media. Conditioned media is harvested and frozen at -20°C until purification.

Conditioned media is purified by affinity chromatography as described below. The media is thawed and then passed through a 0.2 Tm filter. A Protein G column is equilibrated with PBS at pH 7.0, and then loaded with the filtered media. The column is washed with PBS until the absorbance at A₂₈₀ reaches a baseline. IL-1ra-R polypeptide is eluted from the column with 0.1 M Glycine-HCl at pH 2.7 and immediately neutralized with 1 M Tris-HCl at pH 8.5. Fractions containing IL-1ra-R polypeptide are pooled, dialyzed in PBS, and stored at -70°C.

For Factor Xa cleavage of the human IL-1ra-R polypeptide-Fc fusion polypeptide, affinity chromatography-purified protein is dialyzed in 50 mM Tris-HCl, 100 mM NaCl, 2 mM CaCl₂ at pH 8.0. The restriction protease Factor Xa is added to the dialyzed protein at 1/100 (w/w) and the sample digested overnight at room temperature.

10

15

20

25

30

Antibodies to IL-1ra-R polypeptides may be obtained by immunization with purified protein or with IL-1ra-R peptides produced by biological or chemical synthesis. Suitable procedures for generating antibodies include those described in Hudson and Bay, *Practical Immunology* (2nd ed., Blackwell Scientific Publications).

In one procedure for the production of antibodies, animals (typically mice or rabbits) are injected with an IL-1ra-R antigen (such as an IL-1ra-R polypeptide), and those with sufficient serum titer levels as determined by ELISA are selected for hybridoma production. Spleens of immunized animals are collected and prepared as single cell suspensions from which splenocytes are recovered. The splenocytes are fused to mouse myeloma cells (such as Sp2/0-Ag14 cells), are first incubated in DMEM with 200 U/mL penicillin, 200 Fg/mL streptomycin sulfate, and 4 mM glutamine, and are then incubated in HAT selection medium (hypoxanthine, aminopterin, and thymidine). After selection, the tissue culture supernatants are taken from each fusion well and tested for anti-IL-1ra-R antibody production by ELISA.

Alternative procedures for obtaining anti-IL-1ra-R antibodies may also be employed, such as the immunization of transgenic mice harboring human Ig loci for production of human antibodies, and the screening of synthetic antibody libraries, such as those generated by mutagenesis of an antibody variable domain.

Antibodies to IL-1ra-R polypeptides were obtained by immunizing New Zealand White rabbits with full-length IL-1ra-R polypeptide produced in and isolated from *E. coli*. Crude polyclonal immune serum was collected and used in immunoprecipitation analysis using recombinant IL-1ra-R polypeptide (Figures 11A-11B, lanes 1, 4, and 7), recombinant IL-1ra-R polypeptide variant (Figures 11A-11B, lanes 2, 5, and 8), and recombinant IL-1ra (Figures 11A-11B; lanes 3, 6, and 9). Gels were prepared by loading either 10 ng (Figure 11A, lanes 4-6) or 0.6 Π g (Figure 11B, lanes 7-9) of recombinant polypeptide directly onto an 18% Tris-glycine gel or by immunoprecipitating 1 Π g of recombinant polypeptide with 0.2 Π of crude antiserum (Figure 11A, lanes 1-3) prior to loading the sample on

10

15

20

25

30

the gel. Following SDS-PAGE separation, gels were blotted onto PVDF membranes and developed using a 1:1000 dilution of crude anti-IL-1ra-R serum followed by HRP-conjugated Protein A and ECL detection. The gel shown in Figure 11B was stained with Gelcode Blue (Pierce) following blotting. Antibodies to IL-1ra-R polypeptide were shown to detect both membrane-immobilized IL-1ra-R polypeptide and IL-1ra-R polypeptide variant, but not IL-1ra-R polypeptide variant.

Example 7: Expression of IL-1ra-R Polypeptide in Transgenic Mice

To assess the biological activity of IL-1ra-R polypeptide, a construct encoding an IL-1ra-R polypeptide/Fc fusion protein under the control of a liver specific ApoE promoter is prepared. The delivery of this construct is expected to cause pathological changes that are informative as to the function of IL-1ra-R polypeptide. Similarly, a construct containing the full-length IL-1ra-R polypeptide under the control of the beta actin promoter is prepared. The delivery of this construct is expected to result in ubiquitous expression.

To generate these constructs, PCR is used to amplify template DNA sequences encoding an IL-1ra-R polypeptide using primers that correspond to the 5' and 3' ends of the desired sequence and which incorporate restriction enzyme sites to permit insertion of the amplified product into an expression vector. Following amplification, PCR products are gel purified, digested with the appropriate restriction enzymes, and ligated into an expression vector using standard recombinant DNA techniques. For example, amplified IL-1ra-R polypeptide sequences can be cloned into an expression vector under the control of the human E-actin promoter as described by Graham *et al.*, 1997, *Nature Genetics*, 17:272-74 and Ray *et al.*, 1991, *Genes Dev.* 5:2265-73.

Following ligation, reaction mixtures are used to transform an *E. coli* host strain by electroporation and transformants are selected for drug resistance. Plasmid DNA from selected colonies is isolated and subjected to DNA sequencing to confirm the presence of an appropriate insert and absence of

10

15

20

mutation. The IL-1ra-R polypeptide expression vector is purified through two rounds of CsCl density gradient centrifugation, cleaved with a suitable restriction enzyme, and the linearized fragment containing the IL-1ra-R polypeptide transgene is purified by gel electrophoresis. The purified fragment is resuspended in 5 mM Tris, pH 7.4, and 0.2 mM EDTA at a concentration of 2 mg/mL.

Single-cell embryos from BDF1 x BDF1 bred mice are injected as described (PCT Pub. No. WO 97/23614). Embryos are cultured overnight in a CO₂ incubator and 15-20 two-cell embryos are transferred to the oviducts of a pseudopregnant CD1 female mice. Offspring obtained from the implantation of microinjected embryos are screened by PCR amplification of the integrated transgene in genomic DNA samples as follows. Ear pieces are digested in 20 mL ear buffer (20 mM Tris, pH 8.0, 10 mM EDTA, 0.5% SDS, and 500 mg/mL proteinase K) at 55°C overnight. The sample is then diluted with 200 mL of TE, and 2 mL of the ear sample is used in a PCR reaction using appropriate primers.

At 8 weeks of age, transgenic founder animals and control animals are sacrificed for necropsy and pathological analysis. Portions of spleen are removed and total cellular RNA isolated from the spleens using the Total RNA Extraction Kit (Qiagen) and transgene expression determined by RT-PCR. RNA recovered from spleens is converted to cDNA using the SuperScriptTM Preamplification System (Gibco-BRL) as follows. A suitable primer, located in the expression vector sequence and 3' to the IL-1ra-R polypeptide transgene, is used to prime cDNA synthesis from the transgene transcripts. Ten mg of total spleen RNA from transgenic founders and controls is incubated with 1 mM of primer for 10 minutes at 70°C and placed on ice. The reaction is then supplemented with 10 mM Tris-HCl, pH 8.3, 50 mM KCl, 2.5 mM MgCl₂, 10 mM of each dNTP, 0.1 mM DTT, and 200 U of SuperScript II reverse transcriptase. Following incubation for 50 minutes at 42°C, the reaction is stopped by heating for 15 minutes at 72°C and digested with 2U of RNase H for 20 minutes at 37°C. Samples are then amplified by PCR using primers specific for IL-1ra-R polypeptide.

30

10

15

20

25

30

Prior to euthanasia, transgenic animals are weighed, anesthetized by isofluorane and blood drawn by cardiac puncture. The samples are subjected to hematology and serum chemistry analysis. Radiography is performed after terminal exsanguination. Upon gross dissection, major visceral organs are subject to weight analysis.

Following gross dissection, tissues (*i.e.*, liver, spleen, pancreas, stomach, the entire gastrointestinal tract, kidney, reproductive organs, skin and mammary glands, bone, brain, heart, lung, thymus, trachea, esophagus, thyroid, adrenals, urinary bladder, lymph nodes and skeletal muscle) are removed and fixed in 10% buffered Zn-Formalin for histological examination. After fixation, the tissues are processed into paraffin blocks, and 3 mm sections are obtained. All sections are stained with hematoxylin and exosin, and are then subjected to histological analysis.

The spleen, lymph node, and Peyer's patches of both the transgenic and the control mice are subjected to immunohistology analysis with B cell and T cell specific antibodies as follows. The formalin fixed paraffin embedded sections are deparaffinized and hydrated in deionized water. The sections are quenched with 3% hydrogen peroxide, blocked with Protein Block (Lipshaw, Pittsburgh, PA), and incubated in rat monoclonal anti-mouse B220 and CD3 (Harlan, Indianapolis, IN). Antibody binding is detected by biotinylated rabbit anti-rat immunoglobulins and peroxidase conjugated streptavidin (BioGenex, San Ramon, CA) with DAB as a chromagen (BioTek, Santa Barbara, CA). Sections are counterstained with hematoxylin.

After necropsy, MLN and sections of spleen and thymus from transgenic animals and control littermates are removed. Single cell suspensions are prepared by gently grinding the tissues with the flat end of a syringe against the bottom of a 100 mm nylon cell strainer (Becton Dickinson, Franklin Lakes, NJ). Cells are washed twice, counted, and approximately 1 x 10⁶ cells from each tissue are then incubated for 10 minutes with 0.5 Tg CD16/32(FcHII/II) Fc block in a 20 TL volume. Samples are then stained for 30 minutes at 2-8°C in a 100 TL volume of PBS (lacking Ca⁺ and Mg⁺), 0.1% bovine serum albumin, and 0.01% sodium

azide with 0.5 Ig antibody of FITC or PE-conjugated monoclonal antibodies against CD90.2 (Thy-1.2), CD45R (B220), CD11b(Mac-1), Gr-1, CD4, or CD8 (PharMingen, San Diego, CA). Following antibody binding, the cells are washed and then analyzed by flow cytometry on a FACScan (Becton Dickinson).

5

10

15

20

25

Example 9: Functionation of IL-1ra-R Polypeptide

To evaluate whether IL-1ra-R polypeptide affects cellular responses to IL-1, IL-18, or other cytokines, the following experiments were performed. First, murine bone marrow cells were transduced with the IL-1ra-R gene by means of retroviral infection as previously described (Yan *et.al.*,1999, *Exp. Hematol*. 27:1409-17). Transduced bone marrow was then used to transplant lethally irradiated recipient mice. Upon hematopoietic recovery, bone marrow and spleen cells from IL-1ra-R-transduced and control mice (*i.e.*, mice transduced with empty retroviral vector) were subjected to a series of analyses, including FACS analysis (Figure 12), colony assays (Figures 13A-13B), and 9-interferon production in response to either IL-12 (Figure 14) or IL-12 plus IL-18 (Figure 15).

The data suggest that IL-1ra-R gene expression affects the relative abundance and/or cytokine response of specific subsets of hematopoietic progenitors in the spleen (most prominently G-CSF induced GM-CFC; SCF/Epo induced BFU-E/CFU-Mix; and CSF-1 induced M-CFC) and bone marrow (IL-3 and G-CSF induced GM-CFC; SCF/Epo induced BFU-E/CFU-Mix and IL-3/Epo induced GM-CFC/CFU-Mix).

Expression of IL-1ra-R polypeptide also correlates with decreased surface expression of CD4 and NK1.1 on splenocytes and with a decreased ability of splenocytes to produce 9-interferon in response to IL-12 alone or IL-12 plus IL-18.

While the present invention has been described in terms of the preferred embodiments, it is understood that variations and modifications will occur to those skilled in the art. Therefore, it is intended that the appended claims cover

all such equivalent variations that come within the scope of the invention as claimed.

10

15

20

25

30

WHAT IS CLAIMED IS:

- 1. An isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:
- (a) the nucleotide sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35;
- (b) the nucleotide sequence of the DNA insert in ATCC Deposit No. PTA-1423;
- (c) a nucleotide sequence encoding the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
 - (d) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (c); and
 - (e) a nucleotide sequence complementary to any of (a) (c).
 - 2. An isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:
 - (a) a nucleotide sequence encoding a polypeptide which is at least about 70 percent identical to the polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
 - (b) a nucleotide sequence encoding an allelic variant or splice variant of the nucleotide sequence as set forth in any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the nucleotide sequence of the DNA insert in ATCC Deposit No. PTA-1423, or (a);
 - (c) a region of the nucleotide sequence of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the DNA insert in ATCC Deposit No. PTA-1423, (a), or (b) encoding a polypeptide fragment of at least about 25 amino acid residues, wherein the polypeptide fragment has an activity of the encoded polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or is antigenic;

15

20

25

- (d) a region of the nucleotide sequence of any of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, or SEQ ID NO: 35, the DNA insert in ATCC Deposit No. PTA-1423, or any of (a) (c) comprising a fragment of at least about 16 nucleotides;
- (e) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (d); and
 - (f) a nucleotide sequence complementary to any of (a) (d).
- 3. An isolated nucleic acid molecule comprising a nucleotide sequence selected from the group consisting of:
 - (a) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one conservative amino acid substitution, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
 - (b) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid insertion, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
 - (c) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid deletion, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
 - (d) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 which has a C-and/or N- terminal truncation, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;

- (e) a nucleotide sequence encoding a polypeptide as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one modification selected from the group consisting of amino acid substitutions, amino acid insertions, amino acid deletions, C-terminal truncation, and N-terminal truncation, wherein the encoded polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (f) a nucleotide sequence of any of (a) (e) comprising a fragment of at least about 16 nucleotides;
- (g) a nucleotide sequence which hybridizes under moderately or highly stringent conditions to the complement of any of (a) (f); and
 - (h) a nucleotide sequence complementary to any of (a) (e).
- 4. A vector comprising the nucleic acid molecule of any of Claims 1, 2, or 3.
 - 5. A host cell comprising the vector of Claim 4.
 - 6. The host cell of Claim 5 that is a eukaryotic cell.
 - 7. The host cell of Claim 5 that is a prokaryotic cell.
- 8. A process of producing an IL-1ra-R polypeptide comprising culturing the host cell of Claim 5 under suitable conditions to express the polypeptide, and optionally isolating the polypeptide from the culture.
 - 9. A polypeptide produced by the process of Claim 8.
- 10. The process of Claim 8, wherein the nucleic acid molecule comprises promoter DNA other than the promoter DNA for the native IL-1ra-R polypeptide operatively linked to the DNA encoding the IL-1ra-R polypeptide.

10

15

20

25

- 11. The isolated nucleic acid molecule according to Claim 2, wherein the percent identity is determined using a computer program selected from the group consisting of GAP, BLASTN, FASTA, BLASTA, BLASTX, BestFit, and the Smith-Waterman algorithm.
- 12. A process for determining whether a compound inhibits IL-1ra-R polypeptide activity or IL-1ra-R polypeptide production comprising exposing a cell according to any of Claims 5, 6, or 7 to the compound and measuring IL-1ra-R polypeptide activity or IL-1ra-R polypeptide production in said cell.
- 13. An isolated polypeptide comprising the amino acid sequence selected from the group consisting of:
- (a) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36; and
- (b) the amino acid sequence encoded by the DNA insert in ATCC Deposit No. PTA-1423.
- 14. An isolated polypeptide comprising the amino acid sequence selected from the group consisting of:
- (a) an amino acid sequence for an ortholog of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (b) an amino acid sequence which is at least about 70 percent identical to the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (c) a fragment of the amino acid sequence set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 comprising at least about 25 amino acid residues, wherein the fragment has an activity of the

10

15

20

25

30

polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or is antigenic; and

- (d) an amino acid sequence for an allelic variant or splice variant of the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, the amino acid sequence encoded by the DNA insert in ATCC Deposit No. PTA-1423, (a), or (b).
- 15. An isolated polypeptide comprising the amino acid sequence selected from the group consisting of:
- (a) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one conservative amino acid substitution, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (b) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid insertion, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (c) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one amino acid deletion, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36;
- (d) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 which has a C- and/or N- terminal truncation, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36; and
- (e) the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36 with at least one modification selected from the group consisting of amino acid substitutions, amino acid insertions, amino acid deletions, C-terminal truncation, and N-terminal truncation,

20

wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.

- 16. An isolated polypeptide encoded by the nucleic acid molecule of any of Claims 1, 2, or 3, wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.
- 17. The isolated polypeptide according to Claim 14, wherein the percent identity is determined using a computer program selected from the group consisting of GAP, BLASTP, FASTA, BLASTA, BLASTX, BestFit, and the Smith-Waterman algorithm.
 - 18. A selective binding agent or fragment thereof which specifically binds the polypeptide of any of Claims 13, 14, 15, 55, or 56.
 - 19. The selective binding agent or fragment thereof of Claim 18 that specifically binds the polypeptide comprising the amino acid sequence as set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36, or a fragment thereof.
 - 20. The selective binding agent of Claim 18 that is an antibody or fragment thereof.
- 25 21. The selective binding agent of Claim 18 that is a humanized antibody.
 - 22. The selective binding agent of Claim 18 that is a human antibody or fragment thereof.

20

25

- 23. The selective binding agent of Claim 18 that is a polyclonal antibody or fragment thereof.
- 24. The selective binding agent Claim 18 that is a monoclonal antibody or fragment thereof.
 - 25. The selective binding agent of Claim 18 that is a chimeric antibody or fragment thereof.
- 10 26. The selective binding agent of Claim 18 that is a CDR-grafted antibody or fragment thereof.
 - 27. The selective binding agent of Claim 18 that is an antiidiotypic antibody or fragment thereof.

28. The selective binding agent of Claim 18 that is a variable region fragment.

- 29. The variable region fragment of Claim 28 that is a Fab or a Fab' fragment.
 - 30. A selective binding agent or fragment thereof comprising at least one complementarity determining region with specificity for a polypeptide having the amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.
 - 31. The selective binding agent of Claim 18 that is bound to a detectable label.
- 30 32. The selective binding agent of Claim 18 that antagonizes IL-1ra-R polypeptide biological activity.

33. A method for treating, preventing, or ameliorating an IL-1ra-R polypeptide-related disease, condition, or disorder comprising administering to a patient an effective amount of a selective binding agent according to Claim 18.

5

- 34. A selective binding agent produced by immunizing an animal with a polypeptide comprising an amino acid sequence of any of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, or SEQ ID NO: 36.
- 10 35. A hybridoma which produces a selective binding agent which is capable of binding a polypeptide according to any of Claims 1, 2, or 3.
 - 36. A method of detecting or quantitating the amount of IL-1ra-R polypeptide using the anti-IL-1ra-R antibody or fragment of Claim 18.

15

- 37. A composition comprising the polypeptide of any of Claims 13, 14, 15, 55, or 56, and a pharmaceutically acceptable formulation agent.
- 38. The composition of Claim 37, wherein the pharmaceutically acceptable formulation agent is a carrier, adjuvant, solubilizer, stabilizer, or antioxidant.
 - 39. A polypeptide comprising a derivative of the polypeptide of any of Claims 13, 14, 15, 55, or 56.

- 40. The polypeptide of Claim 39 that is covalently modified with a water-soluble polymer.
- The polypeptide of Claim 40, wherein the water-soluble polymer is selected from the group consisting of polyethylene glycol, monomethoxy-polyethylene glycol, dextran, cellulose, poly-(N-vinyl pyrrolidone) polyethylene

20

glycol, propylene glycol homopolymers, polypropylene oxide/ethylene oxide copolymers, polyoxyethylated polyols, and polyvinyl alcohol.

- 42. A composition comprising a nucleic acid molecule of any of Claims 1, 2, or 3 and a pharmaceutically acceptable formulation agent.
 - 43. The composition of Claim 42, wherein said nucleic acid molecule is contained in a viral vector.
- 10 44. A viral vector comprising a nucleic acid molecule of any of Claims 1, 2, or 3.
 - 45. A fusion polypeptide comprising the polypeptide of any of Claims 13, 14, 15, 55, or 56 fused to a heterologous amino acid sequence.
 - 46. The fusion polypeptide of Claim 45, wherein the heterologous amino acid sequence is an IgG constant domain or fragment thereof.
 - 47. A method for treating, preventing, or ameliorating a medical condition comprising administering to a patient the polypeptide of any of Claims 13, 14, 15, 55, or 56, or the polypeptide encoded by the nucleic acid of any of Claims 1, 2, or 3.
- 48. A method of diagnosing a pathological condition or a susceptibility to a pathological condition in a subject comprising:
 - (a) determining the presence or amount of expression of the polypeptide of any of Claims 13, 14, 15, 55, or 56, or the polypeptide encoded by the nucleic acid molecule of any of Claims 1, 2, or 3 in a sample; and
- (b) diagnosing a pathological condition or a susceptibility to a pathological condition based on the presence or amount of expression of the polypeptide.

- 49. A device, comprising:
- (a) a membrane suitable for implantation; and
- (b) cells encapsulated within said membrane, wherein said cells secrete a protein of any of Claims 13, 14, 15, 55, or 56; and

said membrane is permeable to said protein and impermeable to materials detrimental to said cells.

- 50. A method of identifying a compound which binds to an IL-1ra-R polypeptide comprising:
 - (a) contacting the polypeptide of any of Claims 13, 14, 15, 55, or 56 with a compound; and
 - (b) determining the extent of binding of the IL-1ra-R polypeptide to the compound.

15

- 51. The method of Claim 50, further comprising determining the activity of the polypeptide when bound to the compound.
- 52. A method of modulating levels of a polypeptide in an animal comprising administering to the animal the nucleic acid molecule of any of Claims 1, 2, or 3.
 - 53. A transgenic non-human mammal comprising the nucleic acid molecule of any of Claims 1, 2, or 3.

25

54. A process for determining whether a compound inhibits IL-1ra-R polypeptide activity or IL-1ra-R polypeptide production comprising exposing a transgenic mammal according to Claim 53 to the compound, and measuring IL-1ra-R polypeptide activity or IL-1ra-R polypeptide production in said mammal.

10

15

20

25

30

An isolated polypeptide comprising the amino acid sequence as set 55. forth in either SEQ ID NO: 1 or SEQ ID NO: 3 with at least one amino acid substitution selected from the group consisting of: arginine at position 2; alanine, lysine, or arginine at position 3; serine at position 7; lysine at position 8; alanine, cysteine, lysine, threonine, or serine at position 9; cysteine or phenylalanine at position 10; arginine or trptophan at position 13; serine at position 15; arginine at position 18; serine or threonine at position 19; threonine at position 21; serine at position 23; arginine at position 34; tyrosine, serine, or arginine at position 37; lysine, arginine, threonine, or serine at position 38; threonine at position 41; serine, phenylalanine, or alanine at position 43; alanine at position 44; serine or lysine at position 48; alanine, threonine, or phenylalanine at position 52; serine at position 53; serine at position 54; alanine or tyrosine at position 58; lysine at position 65; phenylalanine at position 66; tyrosine at position 67; serine, tyrosine, or phenylalanine at position 69; lysine or serine at position 73; threonine or arginine at position 78; serine or alanine at position 90; alanine at position 91; serine at position 96; lysine or arginine at position 97; lysine or serine at position 98; alanine at position 100; tyrosine at position 102; arginine or alanine at position 104; lysine at position 105; threonine at position 106; arginine at position 108; lysine, threonine, or trptophan at position 109; threonine or serine at position 110; serine at position 111; serine at position 114; serine at position 116; phenylalanine, cysteine, or tyrosine at position 117; tyrosine at position 121; serine or alanine at position 123; cysteine, serine, or threonine at position 126; serine at position 136; phenylalanine or arginine at position 138; threonine, arginine, or alanine at position 141; lysine or tyrosine at position 142; trptophan or threonine at position 143; alanine at position 145; threonine or serine at position 147; cysteine at position 151; and serine, cysteine, or phenylalanine at position 152; wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, or SEQ ID NO: 6.

56. An isolated polypeptide comprising the amino acid sequence as set forth in SEQ ID NO: 5 with at least one amino acid substitution selected from the

5

10

15

20

25

group consisting of: arginine at position 21; alanine, lysine, or arginine at position 22; serine at position 26; lysine at position 27; alanine, cysteine, lysine, threonine, or serine at position 28; cysteine or phenylalanine at position 29; arginine or trptophan at position 32; serine at position 34; arginine at position 37; serine or threonine at position 38; threonine at position 40; serine at position 42; arginine at position 53; tyrosine, serine, or arginine at position 56; lysine, arginine, threonine, or serine at position 57; threonine at position 60; serine, phenylalanine, or alanine at position 62; alanine at position 63; serine or lysine at position 67; alanine, threonine, or phenylalanine at position 71; serine at position 72; serine at position 73; alanine or tyrosine at position 77; lysine at position 84; phenylalanine at position 85; tyrosine at position 86; serine, tyrosine, or phenylalanine at position 88; lysine or serine at position 92; threonine or arginine at position 97; serine or alanine at position 109; alanine at position 110; serine at position 115; lysine or arginine at position 116; lysine or serine at position 117; alanine at position 119; tyrosine at position 121; arginine or alanine at position 123; lysine at position 124; threonine at position 125; arginine at position 127; lysine, threonine, or trptophan at position 128; threonine or serine at position 129; serine at position 130; serine at position 133; serine at position 135; phenylalanine, cysteine, or tyrosine at position 136; tyrosine at position 140; serine or alanine at position 142; cysteine, serine, or threonine at position 145; serine at position 155; phenylalanine or arginine at position 157; threonine, arginine, or alanine at position 160; lysine or tyrosine at position 161; trptophan or threonine at position 162; alanine at position 164; threonine or serine at position 166; cysteine at position 170; and serine, cysteine, or phenylalanine at position 171; wherein the polypeptide has an activity of the polypeptide set forth in any of SEQ ID NO: 2, SEQ ID NO: 4, or SEQ ID NO: 6.

ABSTRACT

The present invention provides novel Interleukin-1 Receptor Antagonist-Related (IL-1ra-R) polypeptides and nucleic acid molecules encoding the same. The invention also provides selective binding agents, vectors, host cells, and methods for producing IL-1ra-R polypeptides. The invention further provides pharmaceutical compositions and methods for the diagnosis, treatment, amelioration, and/or prevention of diseases, disorders, and conditions associated with IL-1ra-R polypeptides.

FIG. 1A

cago	ggato	ag g	gtto	ccago	ja ac	ctcaç	ggato	tgo	agto	jagg	acca	ıgaca	acc a	actga	attgca	60
gga														tat Tyr		108
gac Asp	cag Gln	aag Lys	gct Ala	cta Leu 20	tac Tyr	aca Thr	aga Arg	gat Asp	ggc Gly 25	cag Gln	ctg Leu	ctg Leu	gtg Val	gga Gly 30	gat Asp	156
cct Pro	gtt Val	gca Ala	gac Asp 35	aac Asn	tgc Cys	tgt Cys	gca Ala	gag Glu 40	aag Lys	atc Ile	tgc Cys	aca Thr	ctt Leu 45	cct Pro	aac Asn	204
aga Arg	ggc Gly	ttg Leu 50	gac Asp	cgc Arg	acc Thr	aag Lys	gtc Val 55	ccc Pro	att Ile	ttc Phe	ctg Leu	60 999	atc Ile	cag Gln	gga Gly	252
gly ggg	agc Ser 65	cgc Arg	tgc Cys	ctg Leu	gca Ala	tgt Cys 70	gtg Val	gag Glu	aca Thr	gaa Glu	gag Glu 75	gly ggg	cct Pro	tcc Ser	cta Leu	300
cag Gln 80	ctg Leu	gag Glu	gat Asp	gtg Val	aac Asn 85	att Ile	gag Glu	gaa Glu	ctg Leu	tac Tyr 90	aaa Lys	ggt Gly	ggt Gly	gaa Glu	gag Glu 95	348
														ttc Phe 110		396
														gca Ala		444
														gcc Ala		492
acc Thr	aag Lys 145	ttt Phe	tac Tyr	ttt Phe	gaa Glu	cag Gln 150	agc Ser	tgg Trp	tag	gga	gaca	gga i	aact	gcgt	tt	542
tag	cctt	gtg	cccc	caaa	cc a	agct	catc	c tg	ctca	gggt	cta	tggt	agg	caga	ataatg	602
tcc	cccg	aaa	tatg	tcca	ca t	ccta	atcc	c aa	gatc	tgtg	cat	atgt	tac	cata	catgtc	662
caa	agag	gtt	ttgc	aaat	gt g	atta	tgtt	a ag	gatc	ttga	aat	gagg	aga	caat	cctggg	722
tta	tcct	tgt ·	gggc	tcag	tt t	aatc	acaa	g aa	ggag	gcag	gaa	ggga	gag	tcag	agagag	782
aat	ggaa	gat	acca	tgct	tc t	aatt	ttga	a ga	tgga	gtga	999	gcct	tga	gcca	acaaat	842
gca	ggtg	ttt	ttag	aagg	tg g	aaaa	gcca	a gg	gaac	ggat	tct	cctc	tag	agtc	tccgga	902

FIG. 1B

aggaacacag ctcttgacac atggatttca gctcagtgac acccatttca gacttctgac 962 ctccacaact ataaaataat aaacttgtgt tattgtaaac ctctaaaaaa aaaaaaaa 1020

FIG. 2A

cagg	gato	ag g	gtto	cago	ja ac	ctcag	gato	tgo	cagto	gagg	acca	agaca	acc a	actga	ittgca	60
														tat Tyr		108
gac Asp	cag Gln	aag Lys	gct Ala	cta Leu 20	tac Tyr	aca Thr	aga Arg	gat Asp	ggc Gly 25	cag Gln	ctg Leu	ctg Leu	gtg Val	gga Gly 30	gat Asp	156
cct Pro	gtt Val	gca Ala	gac Asp 35	aac Asn	tgc Cys	tgt Cys	gca Ala	gag Glu 40	aag Lys	atc Ile	tgc Cys	ata Ile	ctt Leu 45	cct Pro	aac Asn	204
aga Arg	ggc Gly	ttg Leu 50	gcc Ala	cgc Arg	acc Thr	aag Lys	gtc Val 55	ccc Pro	att Ile	ttc Phe	ctg Leu	60 Gly 60	atc Ile	cag Gln	gga Gly	252
gly ggg	agc Ser 65	cgc Arg	tgc Cys	ctg Leu	gca Ala	tgt Cys 70	gtg Val	gag Glu	aca Thr	gaa Glu	gag Glu 75	ggg Gly	cct Pro	tcc Ser	cta Leu	300
cag Gln 80	ctg Leu	gag Glu	gat Asp	gtg Val	aac Asn 85	att Ile	gag Glu	gaa Glu	ctg Leu	tac Tyr 90	aaa Lys	ggt Gly	ggt Gly	gaa Glu	gag Glu 95	348
														ttc Phe 110		396
ctt Leu	gag Glu	gct Ala	gct Ala 115	gcc Ala	tgg Trp	cct Pro	ggc Gly	tgg Trp 120	ttc Phe	ctg Leu	tgt Cys	ggc Gly	ccg Pro 125	gca Ala	gag Glu	444
ccc Pro	cag Gln	cag Gln 130	cca Pro	gta Val	cag Gln	ctc Leu	acc Thr 135	aag Lys	gag Glu	agt Ser	gag Glu	ccc Pro 140	tca Ser	gcc Ala	cgt Arg	492
acc Thr	aag Lys 145	ttt Phe	tac Tyr	ttt Phe	gaa Glu	cag Gln 150	agc Ser	tgg Trp	tag	gga	gaca	gga (aact	gcgt [.]	tt	542
tago	cctt	gtg	cccc	caaa	cc a	agct	catc	c tg	ctca	gggt	cta	tggt.	agg	caga	ataatg	602
tccc	cccg	aaa	tatg	tcca	ca t	ccta	atcc	c aa	gatc	tgtg	cat	atgt	tac	cata	catgtc	662
caaa	agag	gtt	ttgc	aaat	gt g	atta	tgtt	a ag	gatc	ttga	aat	gagg	aga	caat	cctggg	722
ttat	cct	tgt (gggc	tcag	tt t	aatc	acaa	g aa	ggag	gcag	gaa	ggga	gag	tcag	agagag	782
aatg	ggaa	gat	acca	tgct	tc t	aatt	ttga	a ga	tgga	gtga	aaa	gcct	tga	gcca	acaaat	842
gcaç	ggtg	ttt	ttag	aagg	tg g	aaaa	gcca	a gg	gaac	ggat	tct	cctc	tag	agtc	tccgga	902

FIG. 2B

aggaacacag ctcttgacac atggatttca gctcagtgac acccatttca gacttctgac 962 ctccacaact ataaaataat aaacttgtgt tattgtaaac ctctaaaaaa aaaaaaaa 1020

FIG. 3

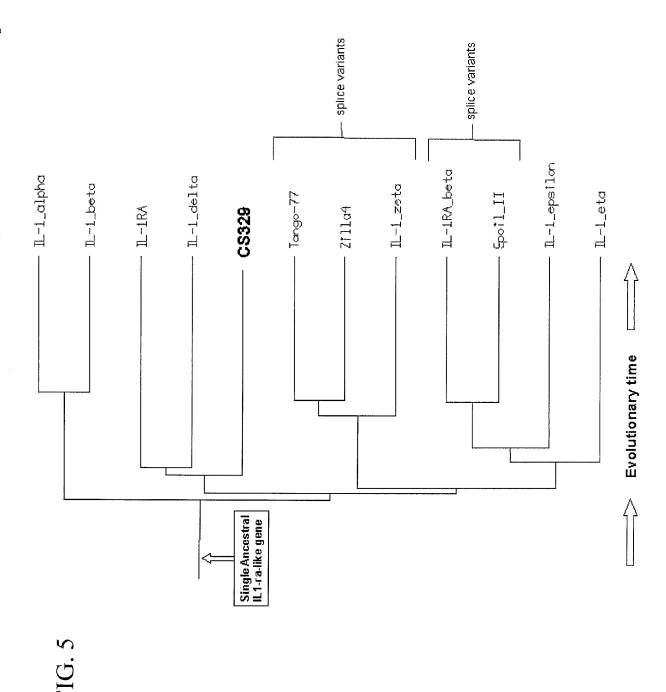
gcto	cccgc	ca g	gaga	aagg	ja ac	atto	tgag	3 999	gagto	ctac	acco	ctgtg	gga g	gctca	ag	57
atg Met 1	gtc Val	ctg Leu	agt Ser	ggg Gly 5	gcg Ala	ctg Leu	tgc Cys	ttc Phe	cgt Arg 10	gag Glu	gac Asp	cag Gln	aca Thr	cca Pro 15	ctg Leu	105
att Ile	gca Ala	gga Gly	atg Met 20	tgt Cys	tcc Ser	ctc Leu	ccc Pro	atg Met 25	gca Ala	aga Arg	tac Tyr	tac Tyr	ata Ile 30	att Ile	aaa Lys	153
tat Tyr	gca Ala	gac Asp 35	cag Gln	aag Lys	gct Ala	cta Leu	tac Tyr 40	aca Thr	aga Arg	gat Asp	ggc Gly	cag Gln 45	ctg Leu	ctg Leu	gtg Val	201
gga Gly	gat Asp 50	cct Pro	gtt Val	gca Ala	gac Asp	aac Asn 55	tgc Cys	tgt Cys	gca Ala	gag Glu	aag Lys 60	atc Ile	tgc Cys	ata Ile	ctt Leu	249
cct Pro 65	aac Asn	aga Arg	ggc Gly	ttg Leu	gcc Ala 70	cgc Arg	acc Thr	aag Lys	gtc Val	ccc Pro 75	att Ile	ttc Phe	ctg Leu	Gly 999	atc Ile 80	297
cag Gln	gga Gly	gly ggg	agc Ser	cgc Arg 85	tgc Cys	ctg Leu	gca Ala	tgt Cys	gtg Val 90	gag Glu	aca Thr	gaa Glu	gag Glu	999 Gly 95	cct Pro	345
tcc Ser	cta Leu	cag Gln	ctg Leu 100	gag Glu	gat Asp	gtg Val	aac Asn	att Ile 105	gag Glu	gaa Glu	ctg Leu	tac Tyr	aaa Lys 110	ggt Gly	ggt Gly	393
gaa Glu	gag Glu	gcc Ala 115	aca Thr	cgc Arg	ttc Phe	acc Thr	ttc Phe 120	ttc Phe	cag Gln	agc Ser	agc Ser	tca Ser 125	ggc Gly	tcc Ser	gcc Ala	441
ttc Phe	agg Arg 130	ctt Leu	gag Glu	gct Ala	gct Ala	gcc Ala 135	tgg Trp	cct Pro	ggc Gly	tgg Trp	ttc Phe 140	ctg Leu	tgt Cys	ggc Gly	ccg Pro	489
gca Ala 145	gag Glu	ccc Pro	cag Gln	cag Gln	cca Pro 150	gta Val	cag Gln	ctc Leu	acc Thr	aag Lys 155	Glu	agt Ser	gag Glu	ccc Pro	tca Ser 160	537
gcc Ala	cgt Arg	acc Thr	aag Lys	ttt Phe 165	Tyr	ttt Phe	gaa Glu	cag Gln	agc Ser 170	Trp	tag	gga	gaca	gga		583
aac	tgcg	ttt	tagc	cttg	tg c	cccc	aaac	c aa	gctc	atco	tgc	tcag	ggt	ctat	ggtagg	643
cag	aata	atg	taca	ccga	aa t	atgt	ccac	a to	ctaa	tccc.	aag	atct	gtg	cata	tgttac	703
cat	acat	qtc	caaa	.gagq	tt t	tgca	aatg	t ga	ttat	gtta	ı a					744

FIG. 4A

	1				50
IL-1_alpha	MAEVPKLASE	MMAYYSGNED	DLFFEADGPK	QMKCSFQDLD	LCPLDGGIQI
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
IL-1RA_beta					
	~~~~~~~				
IL-1_epsilon	~~~~~~~~				
IL-I_eta	~~~~~~~	~~~~~~~	~~~~~~	~~~~~~	~~~~~~
	F-1				100
TI_1 alaha	51 RISDHHYSKG	EDON V GLUMIN	MDVI DVMI UD	CDOTTOTATO	100
	KISDNHISKG				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
	~~~~~~~				
7illa4	~~~~~~~	~~MSFVGENS	GVKMGSEDWE	KDEPOCCI ED	PAGSDI.EDGD
	~~~~~~~				
IL-1RA_beta					
Spoil II	~~~~~~~	MRGTPGDADG	GGRAVYOSSE	SNAVGMGLWR	LRPSALTLSP
IL-1 epsilon					
	~~~~~~~				
-					
	101				150
IL-1_alpha	EPIFFDTWDN	EAYVHDAPVR	SLNCTLRDSQ	QKSLVMSGPY	ELKALHLQGQ
	~~~~~~				
IL-1RA	LLFLFHSETI	CRPSGRKSSK	IQAFRIWDVN	QKTFYLRNN.	QLVAGYLQGP
IL-1_delta	~~~~~~~	~~~~MVLSG	ALCFRMKDSA	LKVLYLHNN.	QLLAGGLHAG
CS329	~~~~~~~	~~~~MCSLPM	ARYYIIKYAD	QKALYTRDG.	QLLVGDPVAD
	SLPTMNFVH.				
	SLPTMNFVH.				
	TKGKNSFKKR				
IL-1RA_beta					
Spoii_ii	VEAPAFSAPL	CTLPFPPVCK	PITGTINDLN	QQVWTLQGQ.	NLVAVPRS
IL-1_epsilon	~~~~~~~	~MEKALKIDI	PQQGSIQDIN	HRVWVLQDQ.	TLIAVPRK
1D-1_eca	~~~~~~	~~MNPQREAA	PKSTAIRDSR	QMVWVLSGN.	SLIAAPLS
	151				200
TI1 almha	DMEQQVVFSM	GE41∪∪Gen	CMDRIDITALG	T MERMIT AT CO	200
TI-1 heta	DMEQQVVFSM		PHDKIDINIA	TYENNIALO	APVADAY 5.1.
	NVNLEEKIDV				
IL-1 delta	KVIKGEEISV	ADNEMI DØGI	DUTUAD	TUGGVMCTSC	VASGUETK
CS329	NC.CAEKICT	L'DNEGI'DELK	VDILIG	TUGGGGCTIVA	VETERC DO
Tango-77	KIFFA	LASSLSSA	AEKGSPILLG	VSKGEFCI.VC	DKDKGUGADG
Zilla4	NYIRPEIFFA	LASSLSSA S	AEKGSPILLG	VSKGEFCLVC	DKDKGOGRDG PYDYGÓSH52
IL-1 zeta	NYIRPEIFFA	LASSLSSA	AEKGSPTLLG	VSKGEFCLVC	DKDKGOGADG 
IL-1RA_beta	DSVTPVTVAV	ITCKYPEALE	OGRGDPTYLG	TONDEMCTIVE	EKAGEO DA
Spoil II	DSVTPVTVAV	ITCKYPEALE	OGRGDPIYLG	IONPEMCTIVE	EKVGEO PT
IL-1_epsilon	DRMSPVTIAL	ISCRHVETLE	KDRGNPTYLG	LNGLNICIMC	AKVGDO PT
IL-1 eta	RSIKPVTLHL	IACRDTEFSD	KEKGNMVYLG	IKGKDLCLFC	AEIOGK PT
_		_			

## FIG. 4B

	201				250
IL-1 alpha	LQLESVDPKN	YPKKKMEK	RFVFNKIEIN	NKLEFESAQF	PNWYISTSQA
IL-1 beta	LQLESVDPKN	YPKKKMEK	RFVFNKIEIN	NKLEFESAQF	PNWYISTSQA
IL-1RA	LQLEAVNITD	LSENRKQDKR	.FAFIRSDSG	PTTSFESAAC	PGWFLCTAME
IL-1_delta	LTLEPVNIME	LYLGAKESKS	.FTFYRRDMG	LTSSFESAAY	PGWFLCTVPE
	LQLEDVNIEE	LYKGGEEATR	.FTFFQSSSG	SAFRLEAAAW	PGWFLCGPAE
Tango-77	LQLKKEKLMK	LAAQKESARR	PFIFYRAQVG	SWNMLESAAH	PGWFICTSCN
Zilla4	LQLKKEKLMK	LAAQKESARR	PFIFYRAQVG	SWNMLESAAH	PGWFICTSCN
IL-1_zeta	LQLKKEKLMK	LAAQKESARR	PFIFYRAQVG	SWNMLESAAH	PGWFICTSCN
IL-1RA_beta	LQLKEQKIMD	LYGQPEPV.K	PFLFYRAKTG	RTSTLESVAF	PDWFIA.SSK
Spoil_II	LQLKEQKIMD	LYGQPEPV.K	PFLFYRAKTG	RTSTLESVAF	PDWFIA.SSK
IL-1 epsilon	LQLKEKDIMD	LYNQPEPV.K	SFLFYHSQSG	RNSTFESVAF	PGWFIAVSSE
IL-1_eta	LQLKEKNIMD	LYVEKKAQ.K	PFLFFHNKEG	STSVFQSVSY	PGWFIATSTT
	251			290	)
	ENMPVFL			~~~~~~	)
				~~~~~~	)
IL-1_beta IL-1RA	ENMPVFL ENMPVFL ADQPVSLTNM	.GGTKGGQDI PDEGVMV	TDFTMQFVSS TKFYFQEDE~	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~)
IL-1_beta IL-1RA	ENMPVFL	.GGTKGGQDI PDEGVMV	TDFTMQFVSS TKFYFQEDE~	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~)
IL-1_beta IL-1RA IL-1_delta	ENMPVFL ENMPVFL ADQPVSLTNM	.GGTKGGQDI PDEGVMV PENGGWNAPI	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~	~~~~~~~~)
IL-1_beta IL-1RA IL-1_delta CS329	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~	~~~~~~~	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK	AEMSPSEVSD	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77 Zilla4	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE CNEPVGVTDK	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK IEFSFQPVCK	AEMSPSEVSD	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77 Zilla4 IL-1_zeta IL-1RA_beta	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE CNEPVGVTDK CNEPVGVTDK CNEPVGVTDK RDQPIILTSE	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH FENRKH FENRKH	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK IEFSFQPVCK IEFSFQPVCK TAFELNIND~	AEMSPSEVSD AEMSPSEVSD AEMSPSEVSD	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77 Zilla4 IL-1_zeta IL-1RA_beta Spoil_II	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE CNEPVGVTDK CNEPVGVTDK CNEPVGVTDK RDQPIILTSE RDQPIILTSE	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH FENRKH LGKSYN LGKSYN	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK IEFSFQPVCK IEFSFQPVCK TAFELNIND~ TAFELNIND~	AEMSPSEVSD AEMSPSEVSD AEMSPSEVSD	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77 Zilla4 IL-1_zeta IL-1RA_beta	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE CNEPVGVTDK CNEPVGVTDK CNEPVGVTDK RDQPIILTSE RDQPIILTSE	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH FENRKH LGKSYN LGKSYN	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK IEFSFQPVCK IEFSFQPVCK TAFELNIND~ TAFELNIND~	AEMSPSEVSD AEMSPSEVSD AEMSPSEVSD	
IL-1_beta IL-1RA IL-1_delta CS329 Tango-77 Zilla4 IL-1_zeta IL-1RA_beta Spoil_II IL-1_epsilon	ENMPVFL ENMPVFL ADQPVSLTNM ADQPVRLTQL PQQPVQLTKE CNEPVGVTDK CNEPVGVTDK CNEPVGVTDK RDQPIILTSE RDQPIILTSE	.GGTKGGQDI PDEGVMV PENGGWNAPI SEPSAR FENRKH FENRKH LGKSYN LGKSYN LGKANT	TDFTMQFVSS TKFYFQEDE~ TDFYFQQCD~ TKFYFEQSW~ IEFSFQPVCK IEFSFQPVCK IEFSFQPVCK TAFELNIND~ TAFELNIND~ TDFGLTMLF~	AEMSPSEVSD AEMSPSEVSD	



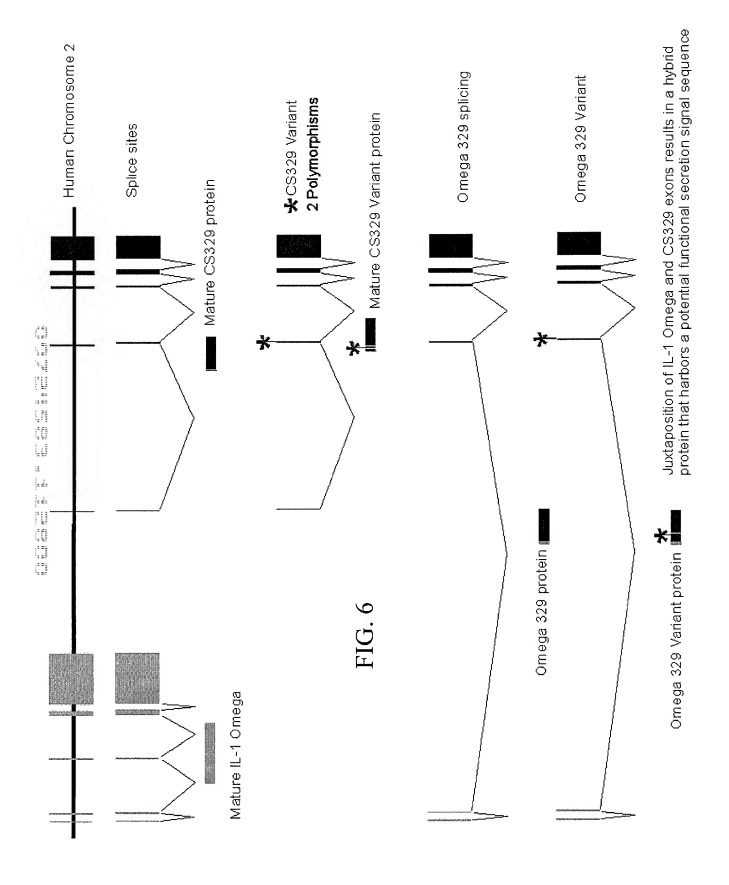


FIG. 7

	tgc Cys															48
	aag Lys															96
	tca Ser															144
ggc Gly	cta Leu 50	gac Asp	cgc Arg	tcc Ser	aag Lys	gtc Val 55	ccc Pro	atc Ile	ttc Phe	ctg Leu	ggg Gly 60	atg Met	cag Gln	gga Gly	gga Gly	192
agt Ser 65	tgc Cys	tgc Cys	ctg Leu	gcg Ala	tgt Cys 70	gta Val	aag Lys	aca Thr	aga Arg	gag Glu 75	gga Gly	cct Pro	ctc Leu	ctg Leu	cag Gln 80	240
ctg Leu	gag Glu	gat Asp	gtg Val	aac Asn 85	atc Ile	gag Glu	gac Asp	cta Leu	tac Tyr 90	aag Lys	gga Gly	ggt Gly	gaa Glu	caa Gln 95	acc Thr	288
	cgt Arg															336
	gct Ala															384
	cag Gln 130															432
_	ttc Phe				_	_		taa								459

FIG. 8

		•
1	${\tt MCSLPMARYYIIKYADQKALYTRDGQLLVGDPVADNCCAEKICTLPNRGL}$	50
1	MCSLPMARYYIIKDAHQKALYTRNGQLLLGDPDSDNYSPEKVCILPNRGL	50
		100
51	DRTKVPIFLGIQGGSRCLACVETEEGPSLQLEDVNIEELYKGGEEATRFT	100
51	DRSKVPIFLGMQGGSCCLACVKTREGPLLQLEDVNIEDLYKGGEQTTRFT	100
101	FFQSSSGSAFRLEAAAWPGWFLCGPAEPQQPVQLTKESEPSARTKFYFEQ	150
		1 - 0
101	FFQRSLGSAFRLEAAACPGWFLCGPAEPQQPVQLTKESEPSTHTEFYFEM	150
a = a	ON 150	
TPT	SW 152	
151	 SR 152	

FIG. 9A

•	actagtctcc	catagacaac	agctgaatgt	acgaggtcag	aagcaaggcc	tgccccagaa	60
,	ccattgcaag	ccaggtgctg	tcttgattgt	agcctcataa	aaaactgatg	cagaattgcc	120
	ccaccaacat	gctccagatt	cctgctccac	agaaaccctg	tgaactaacc	atgttgcttt	180
	tagattctgc	agtaagttga	taatctgcag	taaataacat	tcgatgaaag	agaaacatgt	240
	gtagttactt	tattatgatc	aaaactttat	ttctccactc	tttccatttt	ccttctcaga	300
	attgacacca	gcctttcact	aacccaaata	gcctatttaa	atgctgatca	tacttctctt	360
	gttaactgtt	acctgttccc	aaaaggtaca	attccctttc	gaccatagct	gcatctccca	420
	cctgcacacc	aggatgtttc	tcatatttct	acctaaaaca	ttggggacta	caagtgaaag	480
	caaaagaggg	ggtccatatc	agaaccccag	gtatttagct	gtaaaactca	cttgtcaggc	540
	cagcttgaca	ggtttacagt	ttgtagaagg	accagaaaga	aggtagccaa	gacagaagag	600
	gcaacctctg	cttgtcctag	aaccttcagt	ccatatacat	ctaagctccc	cagcaccatt	660
	tctaccacag	acctctcaga	gttcctgagg	atgcagaccc	caggacactg	acctcagttt	720
	ccaggcaggg	tttctgcaca	ccccttcac	actgcctgac	tgggagttag	tctcatggtg	780
	caacactact	ttgggacact	gtacccatcc	cctcgaccta	cagaaaccat	tcacttttca	840
	aggtcacctc	ctataggaag	tatttgaaaa	gatgagagtc	atggtcattt	gctatgataa	900
	tattctgtgc	ttatctccct	gtaaaaagtt	ggcttggggt	ctctggcatg	catctgacct	960
	taaggttgga	gctgcaccaa	tatgttttta	agcacccggc	ataatgcttc	gcaaaatttc	1020
	agaacatggt	ttgtacagaa	tgtactttcc	tccactcata	caaacccttg	taaaagagta	1080
	gtttgaatcc	caactcattc	ttgaaggcca	ccttttgtag	ggtgacagaa	tttaaaaata	1140
	cagaatttaa	aaatacttta	tcccagggaa	gctcacactt	ctaaatccag	aatgaaagaa	1200
	gaaatagaaa	cacacttgtg	gtggcggtgg	tggtggtgat	ggtggtcgtg	gtggtggtgg	1260
	tggtggtggt	ggtgatggtg	gtggtggtgg	tggtggtggt	ggtcgtggtg	gtgtaatgat	1320
	cacagtaaag	tgaggcatca	tggcctgaga	gagtcaggca	tcacagctat	tcaagtgaaa	1380
	actacctact	actgatttta	gagttctata	attttagtag	cagccacagg	cctggggcct	1440
	gggcctatat	tttcagagag	gaaatgttca	cagcaggtca	actgcagaca	gtgaagatca	1500
	gaaatgtttc	ataatcaggt	catcagagaa	aaggcaaagg	agctgatgga	ctttatcctg	1560
	aaaaagcaaa	atccaaccca	cctcatgctt	aatgcattca	aaggtctgcg	ggcagaagaa	1620

FIG. 9B

tacattttgc	tttttattat	tataaattac	ctggagaata	tttttgtctg	aattatctcc	1680
caaatattaa	ccataaaaat	aaaaaattcc	atgtgtgctt	ctcccagggg	ctataaagcc	1740
cctggtctta	gagttgttgg	ggcaaaacct	gacctttgaa	gtagttactt	ttgaagatgc	1800
cataccatac	atttggccac	ttggagagag	tctaatgtca	catctaaagg	gttactctga	1860
tgctctgttt	tctcatatgc	ccttggctta	cagctaacta	tggctccagc	taaactataa	1920
agttccttgg	caacagagat	ggtacgctat	gtgtctttga	cacagcagaa	taaatgctta	1980
gtgaacatta	ctgattgcct	gacaggacac	ctcacacttt	ggtactttca	acagagggat	2040
gtaaacttat	gaagaacaat	gaagaatgaa	tattggcaat	aaaagcaaaa	attggttaac	2100
ccaattctag	ctctgaaatc	atttttaggt	agtgggaagt	ctttttgttt	tgtttattca	2160
ctttacatcc	caattgctgt	cctccctcca	agttccccac	caccaccaca	gtcctttttc	2220
cctcccttc	tcctctgaga	gaatggagaa	ccctcctgga	tattccccca	tcatgaaaca	2280
ttaagtctct	gcagggctag	acacttcccc	cagtgaggcc	agtcagggca	gcccagctag	2340
aaaaagcata	tcccacagac	agacaacagc	ttttgggata	gccccgttcc	agttgtttag	2400
gatccacatg	aaggctgagc	tgcacatctg	ctacatatga	atgaggaggc	ctaggtccag	2460
cctgtgtatg	ttctttggtt	ggtggttcag	actctgagag	ccccaagggt	ccaggtcagt	2520
tgactctgtt	ggtcttcctg	tggacaccct	gtccccttcc	agcccacaat	ccttccccta	2580
atccttctcc	ttctcacttc	cataagagtg	tgaggagtct	ttaaaaacat	gaagcatttt	2640
atctccccag	ggcaacacat	ggaaatgaaa	gattgtgaaa	agtaatttaa	agaaaaagaa	2700
aaaaaaattt	aacaaggaat	aagaatcttg	tttctctgaa	aatgcttaag	agtgtggaaa	2760
acataaactg	gattctaata	gaatgcaatt	ggattgtaat	gaaaacctat	caaagttatg	2820
aaatagcttt	cactaccttg	cacaaaatct	cttggcatgt	gtgttgttgg	caaattttct	2880
tgttagttta	aaaccacaac	aataacaaca	aaatagcaaa	aattgggtct	cagcctcatt	2940
cattttttct	catttcttgc	tctgtgatcg	tctgggtctt	aagctgacac	ctcaccaatt	3000
cctcatcaag	acctttgtgg	aaatttgcaa	atgtcccaaa	aaggagaatt	acaataagtc	3060
agagaacgtt	ctgtccaatt	ctttatccct	agtgatggat	gagtaaagga	tgtataagag	3120
atggataaat	ggactgatgt	acagataaat	gaaggaatat	gtacatggtt	aggtggatag	3180
atgacttact	caacagatga	gtagaaggat	gagaaataga	tggacagctg	gactgaggca	3240

FIG. 9C

tgcaaagtca	actggagaac	tgagtctctt	gaccatgcac	tgtccagggt	ctcatattcc	3300
ctagagtcca	gggcccatgg	ctcctgtgcc	atccccatgc	aaatctaagg	ttaatacgtt	3360
ctacagctga	gtttccttac	atatgtgtct	cagtaagttt	gtatcaacta	attaaatctg	3420
aaaggagttc	cttctgatct	tcccaaacag	agccacactc	gtgatgaagt	cagccctgct	3480
tcattgtggt	tctctggatg	catctggctt	ccatcagcat	aatctttcta	ttcttgatcc	3540
ttccaacctc	ttcaggtctc	agacagaacc	ccatggagca	tcaaagaggt	ttgaccccag	3600
cattgtttat	gtagctgcaa	aaccactaat	aacacagtca	atgacagtag	ctacagagac	3660
agcaggtcag	tgtctggcct	ctgtcaaggc	tttatgagtg	actctctccc	cttcccgcaa	3720
atactcatta	atctccccac	ctccttatta	tttggactgt	gttgaagata	ttatgaaatc	3780
tctgggctct	tcttcccgga	tctagagcca	attacagatt	ctgtaggttt	gacccaccct	3840
gaccagacat	tataaacaca	gtgctggtgc	cctgaagaaa	acagttggag	actccaggca	3900
ttagaatcca	ggcaccagga	actacaggtc	agtggtgaca	gtcggtctct	ctgtgtatct	3960
cttacacaca	cacacataca	cacacacaac	acaacataca	cacacataca	acacacaaca	4020
catacacata	caacacatac	acacacacaa	cacttttctg	taatgtctcc	aaaattctca	4080
ggctctaggg	aagaagaaat	gtcttttaga	gaatgcggtg	tgatgttcta	taagtctagg	4140
aatacttgat	agaatttaat	gagaagtata	gattaggtca	aagcaagggt	actacatatt	4200
tggaaccaca	gagttttgaa	agtcatctca	aaagaaatta	tttaggccag	agatgttcaa	4260
aaaatgtttt	gtttgtgaca	tatggaagct	cccatggaga	cattctgtga	ttctcatcaa	4320
tagacagtag	ggatgccacc	aaggtgctaa	cgtcttcatc	accccatcat	ctatcataca	4380
tccaaatggt	ttctttgaaa	acaatctcct	tgtgaaactt	aaagtagcct	tgaaaatata	4440
ataatcttgt	ccagcctctc	atttcaatgg	gaatagattg	aaggcctaag	gaccaaaaca	4500
aaaaacaaaa	caaacaaaat	aaaaacacca	aaaaaaaac	ccataaaatg	aatgagtagc	4560
taagttattt	ttagaatcca	gcctttcagt	caaagcttga	ttcatgcata	tctgtgttct	4620
gatcttaagg	tgctgtgtct	gtcagttgta	tagttggata	gaggtacaga	tgagctatat	4680
acatcatgct	tcaagatttc	aggatcttat	aacttttata	aagcaaataa	tttgtcttaa	4740
tgcacactaa	taaacaatat	agcaaagttt	gacaggagtt	cagagtactg	ttagagaagt	4800
gaagggaaga	attttgttat	gatagtaaag	gggaaaatca	aattttgagt	catggaatca	4860

FIG. 9D

tacatagttt gacatagaa	a gaaccttggc	aaccacataa	tctaatgcat	gagcccaaga	4920
actggcctgt gtttttaag	a tctcattctc	agctgttatg	taactgaaca	gacaagatac	4980
taagcccaag tatagtgaa	g ccatgtccag	tgatcttaat	aggagtgaca	ggaatggttg	5040
gtgatgaaga ggggtggat	t ttgagcagga	ataccaaaag	caatgctgac	tgtgcccttg	5100
gagagaatta gcatgagtc	c ttgagagaaa	aatgagatgc	tattgcacaa	gcaacctagg	5160
gccagatggt gtcaagata	g gtggccatcg	tggactttag	aaccaggcag	gaatgtgatc	5220
agagatgtac tttatgtag	g ttaggtttga	ttcagaaacc	aggagggtta	gcatgtttac	5280
aatggtgact aaaaacaag	c acaaggttat	actttaaaga	aataatctct	gaaaagaagg	5340
gaggtatatt ttcagtgcc	g gaaagaggaa	tattacaaaa	gtgagaggag	tagatttgag	5400
aaagagaagt ggattgtgg	a ggagcagatg	ctcaccacgc	ccttacactc	acttgaactg	5460
acacccaaag atgaaggtg	t gctgtggact	gctgaagctc	agcctgtggc	tgggaagcag	5520
taaacaaaat tgctcatca	c agctgtacaa	gatattccat	agcatataaa	aataaaagtg	5580
cttaggctat tctcttaca	a ctctcagcct	tatgaatgac	ccggaaggaa	aagaactcta	5640
caatgtgcct gtgtctgtt	c ttacttcctc	tgccacaagc	aaaagagcct	tgggaattgg	5700
ctcagaggga acgtcatca	a acaggctggc	cttgaggctg	ggctgttatt	cgtctacctg	5760
ggatagagga attcgctat	t cttttataat	ccaagtgtgg	cctggggacc	agcagcatta	5820
ttaagacctg gttgcatgt	t tgaaatgcag	tctcagattt	catcccagac	ctaaagagta	5880
acactgtttt catgaggat	a caagattaag	aaatatgcat	tagagagtaa	ttggctaaat	5940
gggtaaatgt catgcaagc	a ggaggatctg	attgactccc	caggacccac	acagttccca	6000
tgccgtagag cacatctgt	a atcacagtag	gcgtatgatg	aaatgggagg	tgaatcaaga	6060
gaatctctag cagctacgg	g ctggccagcc	tcccatgcac	agcactaaat	aaggcaagga	6120
ccaatacctg aagttgtcc	c attaccttca	catatacacc	acggcatgtg	tgtacttgta	6180
ctcacacata caaacaaat	a cacacgtgca	cacatacaaa	actcagagat	taaggacaat	6240
tggcctgaca tatcagttc	c taagcctggc	tcattgcttg	taacactaca	agcagtatta	6300
aataaggata ggcgagaga	a cagttaccga	atggttcaga	agtggggcca	tgcctgtgac	6360
tttaaacaaa tgtttcata	t ttttaaataa	taacacttag	attacaaaat	aaatttacta	6420
caggaaaatg ttaagaact	a tcaacaacca	ttgactatcc	tgtcggccac	aaatgagtgt	6480

FIG. 9E

tataacaagc	accagccgtc	cttgtccaca	tgtgtgtgtg	tctacacagc	tatgaattta	6540
attgggataa	taatgtgcac	attctttacg	gcctgcagtt	tttacttcat	gtatttgaaa	6600
tgtttgtgcc	acaaatgtca	tctttaagga	gcatatcctt	atttcctgga	tttatcattc	6660
cctttcagcc	gactggacat	tgacagcatt	tccaactttt	caaccttgta	aaaataacta	6720
attgaactat	tttataacta	agcatttggg	caatcaatta	cctctgcctg	gaatgggggc	6780
aacaacacat	gcaatcatgg	gaaagccagg	atgctgctgt	ctgatcccta	gccctggcat	6840
tcgtgcagaa	cctcactctc	atctgtgccc	tgatatcctt	cactctcaag	tcttttccca	6900
gtgactttta	aaggcaacag	aatcatatag	ccaataatga	aagctacttg	gtctacagtt	6960
gtgtggcgtt	ttttatagat	attttcttca	tttacatttc	aaatgctatc	ccaaaagtcc	7020
cctataccct	ccccaccct	gctcccctac	ccactcactc	ccacttcttg	gccctggctt	7080
tcccccttac	tggggcatat	aaagtttgct	agaccaaggg	gcctctcttc	ccaatgatgg	7140
ccaactaggc	cattttctgc	tacatatgca	gctagagaca	ccagttctgg	ggttactggt	7200
tagttcatat	tgttgttcta	cctatggggt	tgcagacccc	ttcagctctt	gagtactttc	7260
ctageteet	ccattgggag	ccctgtgttc	catcctatag	atgactgtga	gcatccactt	7320
ctgtatttgc	caggcactgg	catatgaaat	agtatctgca	tttggtggct	gattatggga	7380
ggacccccg	ggtggggcag	tctctggatg	gtccatcctt	tcatcttagc	tccaaacttt	7440
gtctctgcaa	cttcttccat	ggatatttta	gtccctaatc	tagggagaaa	tgaagtatcc	7500
acaagttgat	cttccttctt	gattttctta	tgttttagaa	gttgtatctt	ggatattcta	7560
ggtttctggg	ctaatatcca	cttatcagtg	agtacatatc	aagtgaattc	ttttgtgatt	7620
aggttacctc	actcaagatg	atattctcca	ctatgttcat	agcagcccta	tttatagtag	7680
ccagaagctg	gaaagaaccc	agtccctcaa	cagaggaatg	gatacagaaa	atgtggcaca	7740
ttatgcaat	ggagtaccac	tcagatatta	aaaacaacga	atttatgaaa	ttctcgggca	7800
aaccctatc	taaagaccag	gaataaggaa	aagatggact	gcctgcctgc	agctgggaga	7860
gctggggaga	cctttgtgga	ttctgtaata	cttaggggta	cggaacagct	tgtggctgga	7920
aattctgag	ctccagcatg	tctgcccccc	aaaaaacatt	ctgtttttct	gaaagccttt	7980
tcttctttg	cctcagtgaa	gaccagacac	tcccaactgc	agga <u>atg</u> to	ge tee ett	8036
cc atg gca	aga tac ta	<u>ıc at</u> gtaagt	aa tottaaco	gat cgctcaat	ca	8084

FIG. 9F

aggggcctgg	agatcacatg	agaagggaaa	aggctgagtc	aaagggacaa	agctccctct	8144
agccacagaa	atctcaaaca	ctgaataatt	gatcttcatc	tttgtcaatc	acaacagccc	8204
tctttcctgg	tgacagaatg	gaacaactgt	aagagtggta	ttgcttagtc	cattttacag	8264
acccggaaac	tcaacctcca	cgaggttata	caattttcct	catgtcatgc	aattacccaa	8324
aagcagagag	tgggatcgga	ctctctgttc	tctaaactga	tgtagctagt	tcttagaaag	8384
ctcaaacaat	cttgagtccc	aaggacagca	cctttatggt	cacctggatt	gatacctata	8444
tcaaaaaaaa	aaaaaggtct	cactagatag	ccctggctac	cctgaaactc	tcactgtgta	8504
catttaggtg	accacgaact	cacagagatc	tgccttccaa	gtgctgggat	taaagtatgt	8564
accaccacac	ctgcatcttt	gacaataact	gagtggtatc	taaattcttc	cagtggctaa	8624
acagttaagt	cccagttccc	aaagtctgag	aaaaatgcca	ggtggtgaaa	tctgtacaga	8684
cctttgttct	taatgtacaa	gtgagcctgc	tttaaaaaca	atacgcaagc	tgtttttgct	8744
attgctaagt	gttgcagaga	cagaaaaggc	tcccagaagt	ggtaactttg	gtccagaggt	8804
tctgttctca	aactcattgt	gagctctgaa	agcaactgat	gggcagctct	gaaatcagct	8864
gggcaattag	gctaataaca	ggcataattt	taatgtttca	cacgcatgac	agttcctccc	8924
cagctgccct	agtacatact	taccctccta	ggcacgtcat	tagacccata	ggtataacca	8984
gtgactaatc	aggccctggt	ctaattctaa	gttggcctcc	tatataagtg	ccactcagag	9044
tgtacctcat	catggctgta	gtgggcccag	agtctaggga	catagacttt	tctattgtcc	9104
aatttctgat	ttgtgaattt	tctacaaaaa	gaatttttt	taattttaca	aatcaaatca	9164
cagttactac	atcttcagtt	ccttcattaa	ttagtgttac	tatttaaaaa	aataaaataa	9224
atcaagctca	gaaacatcat	ggatagggtt	cattgtatct	ccagggtacc	tgagcttcaa	9284
agcaactcct	cagacagcca	tgaaaacatc	ctcaattacc	tcatgagaag	acactattgt	9344
catttctgga	gcctctgata	atcctgagcc	taggcagctt	tgggatgaaa	caatttctac	9404
ccttattgga	acagtgtccc	tctcctgtct	ggaaacaatt	caccaaaggc	tccatgtggt	9464
tgtccagtaa	ggtggtatgg	ggacagaaat	ggacaatgat	ccctgagggc	agtgatccat	9524
taaccttgcc	ctcctatttc	ag <u>a atc aa</u>	ng gat gca c	cat caa aag	gct ttg tac	9577
<u>aca cgg aat</u>	ggc cag ct	c ctg ctg c	ga gac cct	gat tca gad	aat tat	9625
agt cca q	gtgatcttc co	atgataga ac	rtqqqqqaq to	rgagggag gg	rtatagaga	9681

FIG. 9G

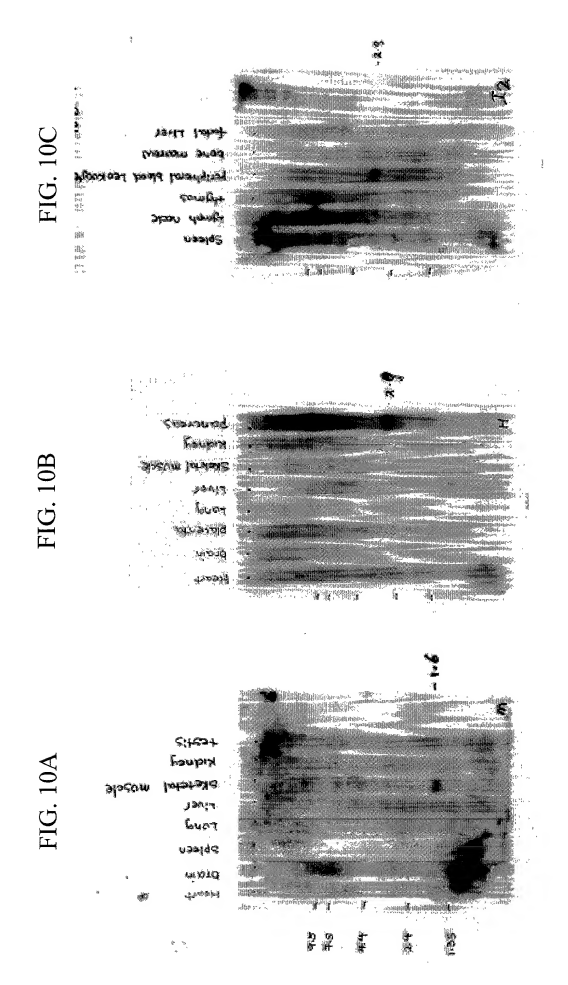
gggctctctt ccagaagttg cttagtgtcc atctgccaca aggccttgat tctttccttc	9741
aattgtgtct ctagagacat gagaatattg tcacagtgat aaggagaaga ggtaggggca	9801
gtttcttcct gtaaaaaatg aattccattt accctgcagt ctccatacag aaacaggcca	9861
gaggggggca gacccagtaa cttctagctg agccctacct tgcttaaaac ctgccatctg	9921
tggtcccctc actgtctgaa ttgcattctg tcttacctcc cag ag aag gtc tgt atc	<u>:</u> 9978
ctt cct aac cga ggc cta gac cgc tcc aag gtc ccc atc ttc ctg ggg	10026
atg cag gga gga agt tgc tgc ctg gcg tgt gta aag aca aga gag gga	10074
cct ctc ctg cag ctg gag gtgagacacc cctcctcatt gcagtcagta	10122
ctgccactgg aacatagtga catctttgaa cccacatgtc ccctctcttg tttcccatct	10182
atctctcttt gcctccagct gagggactct agcctttggg gatgtacaga aagaacatgg	10242
cttcggaaaa ctcttcccta ttgagtcctt ctttggccaa gcctctgagg cactaagggc	10302
tgacgtccca accaaacact catttcatct cacagctgtc tccctttccc cacag gat	10360
gtg aac atc gag gac cta tac aag gga ggt gaa caa acc acc cgt ttc	10408
acc ttt ttc cag aga agc ttg gga tct gcc ttc agg ctt gag gct gct	10456
gcc tgc cct ggc tgg ttt ctc tgt ggc cca gct gag ccc cag cag cca	10504
gtg cag ctc acc aaa gag agt gaa ccc tcc acc cat act gaa ttc tac	10552
ttt gag atg agt cgg taa agagacataa ggctggggcc tcgtctagtg	10600
ccccagtct gagatcttct tgctcagcat ctctggaaag cagaataagg aagataccaa	10660
agatgtttgg gtcttaatcc ccagaatctg tgaccgtgtt acattaaatg gcaaagggat	10720
tttttttttc cttcatggtc catttgggcc cattggaatc atctgaggcc tcatgaggag	10780
aaggaagagg tcagagggag actggggcaa actttggtac taaaagtaac aatggagaca	10840
gggaccataa gctgatgggt aacagtggtt tctagaaacc ggaaatgatg agagctctcc	10900
tgacacaggt tctggatttt tctggactga agaatggtga aataatacag ctccattatt	10960
ttaagccact gagtttgaga tcattcaatg aagctgtcat aataaaacct gtgcttcaca	11020
tacaattcaa tattggtagg cacccggtg atttcttgga aagacatcta gggattctcc	11080
tggatgctga ttccagggtc cagtggagtc cctgggttga agagatttca caacccagaa	11140
catcaggete gactetteta aaagteegte gttgcaceee ttgcctgaga gcattagcaa	11200

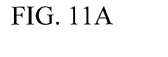
FIG. 9H

tttctatttc	ataggaaatc	tgtgtccctg	cccctgctaa	agcagggagc	ctggaccgtc	11260
ctgatttagt	gaggggtgag	ctgctggcac	ttttttgtgt	caccagtgtc	ttaagcagtg	11320
atggagcaca	aaagatcttt	actgagaaga	tggccatgaa	gctctggcta	gacaccaaga	11380
atatgatata	agcagagcta	cagcacaaga	tgagccaatg	aggaaagcca	ttcagggagg	11440
ctaagcccag	cttcccaaag	ggacagctaa	ccctggactc	aaatgaatag	gggttttcct	11500
ggcagagaac	ataggtcaag	cattctaggt	agaatcagca	attcagaaag	gtgtgagaga	11560
ggcatggaga	gctccaggca	tgtctgggct	atggtgtgtc	attcttgtgg	caagaatcca	11620
acgtctgtgg	ttaaggagtt	gctgaaaatt	aaaataggaa	aatgggtaga	gtctaattgt	11680
gaatgacttg	caaaggagtt	tagcccataa	gtggggagct	cagaggagtc	atctaaggat	11740
tgcaagcagg	ggccctgtga	tcattgctgg	accagcctag	gtgctacaga	gcctaccttc	11800
agctctgcat	cctcactcac	atccaggtac	cttcagaggt	caatttctgt	gctctggttc	11860
tatgggtagc	ctgaccctgt	ttcatcttct	tgtataactt	aggcacataa	gcttagggac	11920
tggtagagtt	tacttgagtg	attggtgaat	caggcagcac	caaactacaa	gttgttcagg	11980
gctttaccaa	gggggcactg	attggagaat	tggaatgagg	gtggttagaa	tgcattcaga	12040
aaacaagggg	aagaaaaatt	tgattgctta	aagtggaaag	tcccaactta	aatgttagtc	12100
agtagtttct	aattacttga	gtctctaatt	agaggttagt	tggcagtttc	tggttagtta	12160
atctaagttt	cattttctta	ggctatgacc	attctctgag	tcgcatgtta	gcaatgcagt	12220
aagaactcaa	gacccagaat	agcctctgtt	aattatttta	gcaatgatca	ctcatttctg	12280
ttgcctccta	ttgagatctg	ttcccatgga	ccacccaggc	acatcaggcc	tcctagtacc	12340
aacataataa	tgattgctgc	acagacaaaa	tattttttt	cagtatctgg	tatttgctac	12400
atttccatta	gtgctggagg	gaaggctaca	acgaccatga	aggcatggcc	cctgccttct	12460
aaggacttac	aatgtaatag	gagccctgac	attataaagt	gggtcacctt	gtttcaaact	12520
gagccaaact	gaggctgagg	gcttagatta	gtggtaggtc	actttccaga	catgttcagt	12580
gctaagaaaa	acacattctg	gggttagtta	gatgttttag	ttcatttgat	aagaagccca	12640
atgattggac	tttcaacttc	tggaacccat	gtggtggaag	agagaaccaa	cttctgacca	12700
tttgggtcat	ggcacatccc	ctaccatcac	aagaactcac	caaaataaat	tagaaaaatc	12760
aagaaaaact	catatcctat	agacctctgg	tagaattagc	agaacgctgc	tgtggcactt	12820

FIG. 9I

gggatttgaa	actcaaaaat	ggaagaagct	acttgtgacc	gttcaagact	ccagggaggc	12880
tcctctgaca	catcccacga	ctcaggctta	aattccttct	tctccctaga	aggccacgcc	12940
atcttctcaa	ccaggccaca	gatgctataa	ttatgtaaat	gtgtgggaga	ggcacacttt	13000
agatcttatc	cactagt					13017





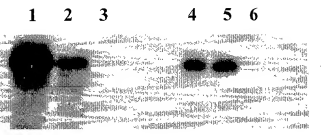


FIG. 11B

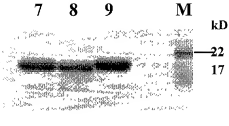


FIG. 12

Spleen

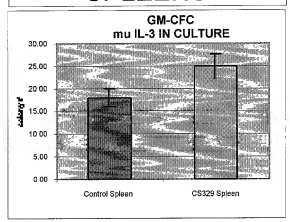
	CD4+	CD8+	CD4+ CD8+	CD3+	NK1.1+	CD3+ NK1.1+
control mice	20.18	3.72	1.67	24.07	3.06	1.4
CS329 mice	15.89	3.99	0.37	22.9	2.08	1.1
Difference:	-4.29	0.27	-1.3	-1.17	86:0-	-0.3

Bone Marrow

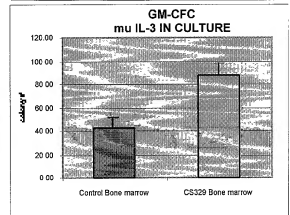
control mice	2.62	CD8+	CD4+CD8+	3.88	NK4.1+	CD34/NKTT#
CS329 mice	2.46	2.35	0.41	4.42	1.53	0.57
35	0.16	0.19	0.08	0.54	0.27	0.08

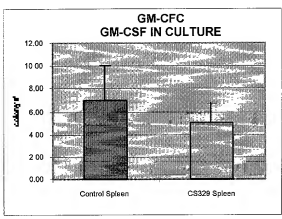
FIG. 13A

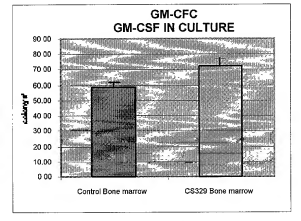
SPLEENS

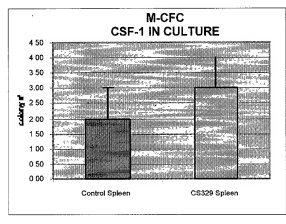


BONE MARROW









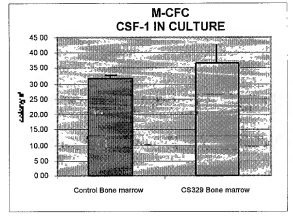
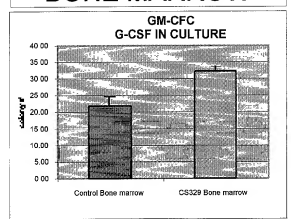


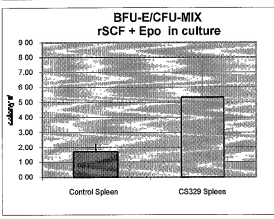
FIG. 13B

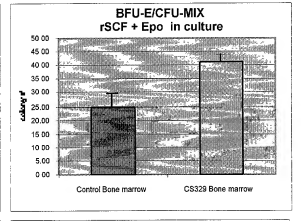
SPLEENS

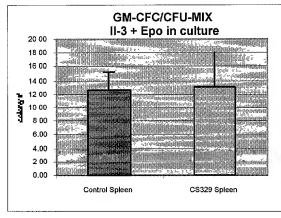
GM-CFC G-CSF IN CULTURE 5 00 4 50 4 00 3 3.00 2 50 2 00 1 50 1 00 0 50 0 00 Control Spleen CS329 Spleen

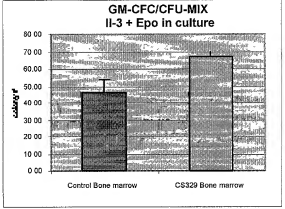
BONE MARROW











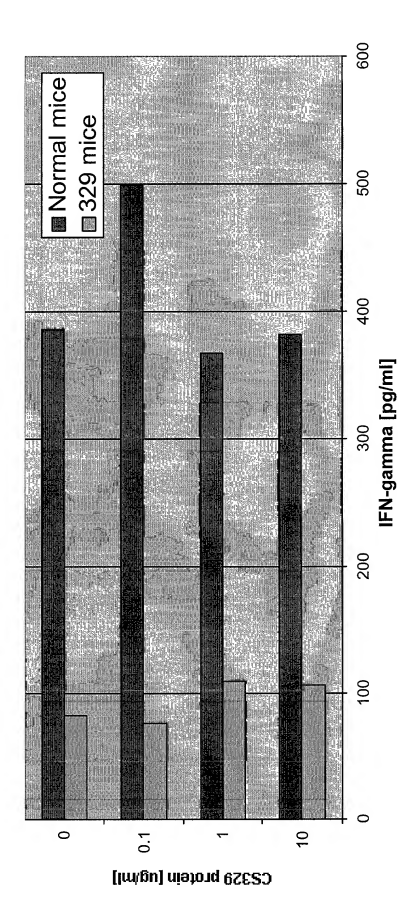
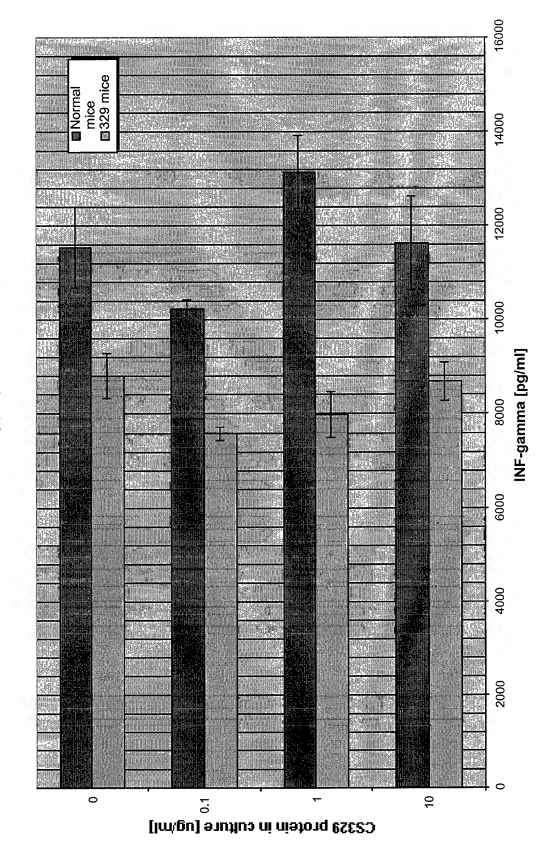


FIG. 15



IN THE UNITED STATES PATENT AND TRADEMARK OFFICE
(Case No. 00,1213)

· ·	PATENT
In re Application of: Welcher et al.)
Serial No.: Unassigned) Before the Examiner: Unassigned
Filed: November 28, 2000) Group Art Unit: Unassigned)
For: Interleukin-1)
Receptor Antagonist-Related)
Molecules and Uses Thereof)

Assistant Commissioner for Patents Washington, D.C. 20231

Sir/Madam:

STATEMENT UNDER 37 C.F.R. § 1.821

The undersigned representative hereby declares that the content of the paper and computer readable copies of the Sequence Listing, submitted in the above-identified application in accordance with 37 C.F.R. §§ 1.821(c) and (e), respectively, are identical in content. The 3.5" diskette contains an IBM compatible dos-text file of the sequence listing named "001213seq.txt."

Respectfully submitted,

McDonnell Boehnen Hulbert & Berghoff

Dated: November 27, 2000

Kevin E. Noonan, Ph.D.

Reg. No. 35,303

SEOUENCE LISTING

```
<110> Saris, Christiaan M.
         Giles, Jennifer
         Mu, Sharon X.
        Xia, Min
        Bass, Michael B.
         Craveiro, Roger
   <120> Interleukin-1 Receptor Antagonist-Related Molecules and
         Uses Thereof
   <130> 00-1213
   <140>
   <141>
   <150> 60/170,191
   <151> 1999-12-10
   <150> 60/188,053
   <151> 2000-03-09
   <150> 60/194,521
   <151> 2000-04-04
<150> 60/195,910
   <151> 2000-04-10
   <160> 32
<170> PatentIn Ver. 2.0
. 4:
   <210> 1
   <211> 1020
   <212> DNA
   <213> Homo sapiens
   <220>
   <221> CDS
   <222> (64)..(522)
   <400> 1
   cagggatcag ggttccagga actcaggatc tgcagtgagg accagacacc actgattgca 60
   gga atg tgt tcc ctc ccc atg gca aga tac tac ata att aaa tat gca
                                                                       108
       Met Cys Ser Leu Pro Met Ala Arg Tyr Tyr Ile Ile Lys Tyr Ala
                                             10
         1
   gac cag aag gct cta tac aca aga gat ggc cag ctg ctg gtg gga gat
                                                                       156
   Asp Gln Lys Ala Leu Tyr Thr Arg Asp Gly Gln Leu Leu Val Gly Asp
                                                              30
                                         25
                     20
   cct gtt gca gac aac tgc tgt gca gag aag atc tgc aca ctt cct aac
                                                                       204
   Pro Val Ala Asp Asn Cys Cys Ala Glu Lys Ile Cys Thr Leu Pro Asn
                                                          45
                 35
```

iŝ

<213> Homo sapiens

20

<400> 2

aga ggc ttg gac cgc acc aag gtc ccc att ttc ctg ggg atc cag gga Arg Gly Leu Asp Arg Thr Lys Val Pro Ile Phe Leu Gly Ile Gln Gly

300

348

396

444

492

542

1020

2

30

10

Met Cys Ser Leu Pro Met Ala Arg Tyr Tyr Ile Ile Lys Tyr Ala Asp

Gln Lys Ala Leu Tyr Thr Arg Asp Gly Gln Leu Leu Val Gly Asp Pro

```
Val Ala Asp Asn Cys Cys Ala Glu Lys Ile Cys Thr Leu Pro Asn Arg
           35
  Gly Leu Asp Arg Thr Lys Val Pro Ile Phe Leu Gly Ile Gln Gly Gly
  Ser Arg Cys Leu Ala Cys Val Glu Thr Glu Glu Gly Pro Ser Leu Gln
                        70
  Leu Glu Asp Val Asn Ile Glu Glu Leu Tyr Lys Gly Gly Glu Glu Ala
  Thr Arg Phe Thr Phe Phe Gln Ser Ser Ser Gly Ser Ala Phe Arg Leu
                                   105
  Glu Ala Ala Trp Pro Gly Trp Phe Leu Cys Gly Pro Ala Glu Pro
                               120
          115
  Gln Gln Pro Val Gln Leu Thr Lys Glu Ser Glu Pro Ser Ala Arg Thr
                           135
  Lys Phe Tyr Phe Glu Gln Ser Trp
                       150
  <210> 3
  <211> 1020
  <212> DNA
  <213> Homo sapiens
  <220>
... <221> CDS
  <222> (64)..(522)
   <400> 3
   cagggatcag ggttccagga actcaggatc tgcagtgagg accagacacc actgattgca 60
   gga atg tgt tcc ctc ccc atg gca aga tac tac ata att aaa tat gca
                                                                      108
       Met Cys Ser Leu Pro Met Ala Arg Tyr Tyr Ile Ile Lys Tyr Ala
   gac cag aag gct cta tac aca aga gat ggc cag ctg ctg gtg gga gat
                                                                      156
   Asp Gln Lys Ala Leu Tyr Thr Arg Asp Gly Gln Leu Leu Val Gly Asp
                    20
   cct gtt gca gac aac tgc tgt gca gag aag atc tgc ata ctt cct aac
                                                                      204
   Pro Val Ala Asp Asn Cys Cys Ala Glu Lys Ile Cys Ile Leu Pro Asn
                35
                                     40
   aga ggc ttg gcc cgc acc aag gtc ccc att ttc ctg ggg atc cag gga
                                                                      252
   Arg Gly Leu Ala Arg Thr Lys Val Pro Ile Phe Leu Gly Ile Gln Gly
                                55
            50
                                                                      300
   ggg agc cgc tgc ctg gca tgt gtg gag aca gaa gag ggg cct tcc cta
   Gly Ser Arg Cys Leu Ala Cys Val Glu Thr Glu Glu Gly Pro Ser Leu
        65
```

7,4

L

cag ctg gag gat gtg aac att gag gaa ctg tac aaa ggt ggt gaa gag Gln Leu Glu Asp Val Asn Ile Glu Glu Leu Tyr Lys Gly Gly Glu Glu 80 85 90 95	340										
gcc aca cgc ttc acc ttc ttc cag agc agc tca ggc tcc gcc ttc agg Ala Thr Arg Phe Thr Phe Phe Gln Ser Ser Ser Gly Ser Ala Phe Arg 100 105 110	396										
ctt gag gct gct gcc tgg cct ggc tgg ttc ctg tgt ggc ccg gca gag Leu Glu Ala Ala Ala Trp Pro Gly Trp Phe Leu Cys Gly Pro Ala Glu 115 120 125	444										
ccc cag cag cca gta cag ctc acc aag gag agt gag ccc tca gcc cgt Pro Gln Gln Pro Val Gln Leu Thr Lys Glu Ser Glu Pro Ser Ala Arg 130 135 140	492										
acc aag ttt tac ttt gaa cag agc tgg tag ggagacagga aactgcgttt Thr Lys Phe Tyr Phe Glu Gln Ser Trp 145 150	542										
tagcettgtg cccccaaacc aagetcatee tgctcagggt ctatggtagg cagaataatg	602										
toccoogaaa tatgtocaca toctaatooo aagatotgtg catatgttac catacatgto	662										
caaagaggtt ttgcaaatgt gattatgtta aggatcttga aatgaggaga caatcctggg	722										
ttatccttgt gggctcagtt taatcacaag aaggaggcag gaagggagag tcagagagag	782										
aatggaagat accatgcttc taattttgaa gatggagtga ggggccttga gccaacaaat	842										
gcaggtgttt ttagaaggtg gaaaagccaa gggaacggat tctcctctag agtctccgga	a 902										
aggaacacag ctcttgacac atggatttca gctcagtgac acccatttca gacttctgac	962										
ctccacaact ataaaataat aaacttgtgt tattgtaaac ctctaaaaaa aaaaaaaa	1020										
<210> 4 <211> 152 <212> PRT <213> Homo sapiens											
<400> 4 Met Cys Ser Leu Pro Met Ala Arg Tyr Tyr Ile Ile Lys Tyr Ala Asp											
1 5 10 15											
Gln Lys Ala Leu Tyr Thr Arg Asp Gly Gln Leu Leu Val Gly Asp Pro 20 25 30											
Val Ala Asp Asn Cys Cys Ala Glu Lys Ile Cys Ile Leu Pro Asn Arg 35 40 45											
Gly Leu Ala Arg Thr Lys Val Pro Ile Phe Leu Gly Ile Gln Gly Gly 50 55 60											
Ser Arg Cys Leu Ala Cys Val Glu Thr Glu Glu Gly Pro Ser Leu Gln 65 70 75 80											

	Leu	Glu	Asp	Val	Asn 85	Ile	Glu	Glu	Leu	Tyr 90	Lys	Gly	Gly	Glu	Glu 95	Ala	
	Thr	Arg	Phe	Thr 100	Phe	Phe	Gln	Ser	Ser 105	Ser	Gly	Ser	Ala	Phe 110	Arg	Leu	
	Glu	Ala	Ala 115	Ala	Trp	Pro	Gly	Trp 120	Phe	Leu	Cys	Gly	Pro 125	Ala	Glu	Pro	
	Gln	Gln 130	Pro	Val	Gln	Leu	Thr 135	Lys	Glu	Ser	Glu	Pro 140	Ser	Ala	Arg	Thr	
	Lys 145	Phe	Tyr	Phe	Glu	Gln 150	Ser	Trp									
	<212	L> 74 2> Di	JA	sapie	ens												
# 1	<220 <220 <222	l> CI		. (573	3)												
## ## ## ## ## ## ## ## ## ## ## ## ##	<400 gct)> 5 cccg	cca 🤉	ggaga	aaag	ga ao	catto	ctga	3 33	gagto	ctac	acco	ctgt	gga g	gctca	aag	57
	atg Met 1	gtc Val	ctg Leu	agt Ser	ggg Gly 5	gcg Ala	ctg Leu	tgc Cys	ttc Phe	cgt Arg 10	gag Glu	gac Asp	cag Gln	aca Thr	cca Pro 15	ctg Leu	105
	att Ile	gca Ala	gga Gly	atg Met 20	tgt Cys	tcc Ser	ctc Leu	ccc Pro	atg Met 25	gca Ala	aga Arg	tac Tyr	tac Tyr	ata Ile 30	att Ile	aaa Lys	153
12 25	tat Tyr	gca Ala	gac Asp 35	cag Gln	aag Lys	gct Ala	cta Leu	tac Tyr 40	aca Thr	aga Arg	gat Asp	ggc Gly	cag Gln 45	ctg Leu	ctg Leu	gtg Val	201
	gga Gly	gat Asp 50	Pro	gtt Val	gca Ala	gac Asp	aac Asn 55	tgc Cys	tgt Cys	gca Ala	gag Glu	aag Lys 60	atc Ile	tgc Cys	ata Ile	ctt Leu	249
	cct Pro 65	aac Asn	aga Arg	ggc	ttg Leu	gcc Ala 70	cgc Arg	acc Thr	aag Lys	gtc Val	ccc Pro 75	att Ile	ttc Phe	ctg Leu	gly	atc Ile 80	297
	cag Gln	gga Gly	Gly 999	agc Ser	cgc Arg 85	tgc Cys	ctg Leu	gca Ala	tgt Cys	gtg Val 90	gag Glu	aca Thr	gaa Glu	gag Glu	95 999	cct Pro	345
	tcc Ser	cta Leu	cag Gln	ctg Leu 100	gag Glu	gat Asp	gtg Val	aac Asn	att Ile 105	Glu	gaa Glu	ctg Leu	tac Tyr	aaa Lys 110	Gly	ggt Gly	393

	gaa Glu	gag Glu	gcc Ala 115	aca Thr	cgc Arg	ttc Phe	acc Thr	ttc Phe 120	ttc Phe	cag Gln	agc Ser	agc Ser	tca Ser 125	ggc Gly	tcc Ser	gcc Ala	441
	ttc Phe	agg Arg 130	ctt Leu	gag Glu	gct Ala	gct Ala	gcc Ala 135	tgg Trp	cct Pro	ggc Gly	tgg Trp	ttc Phe 140	ctg Leu	tgt Cys	ggc Gly	ccg Pro	489
	gca Ala 145	gag Glu	ccc Pro	cag Gln	cag Gln	cca Pro 150	gta Val	cag Gln	ctc Leu	acc Thr	aag Lys 155	gag Glu	agt Ser	gag Glu	ccc Pro	tca Ser 160	537
	gcc Ala	cgt Arg	acc Thr	aag Lys	ttt Phe 165	tac Tyr	ttt Phe	gaa Glu	cag Gln	agc Ser 170	tgg Trp	tag	ggag	gacag	gga		583
	aact	gcgt	ett t	agco	cttgt	g co	ccca	aaac	c aaq	gctca	atcc	tgct	cag	ggt (ctate	ggtagg	643
	caga	ataa	atg t	cccc	ccgaa	aa ta	atgto	ccaca	a tc	ctaat	ccc	aaga	atct	gtg (cata	tgttac	703
1 2 2	cata	acat	gtc (caaag	gaggt	t tt	gcaa	aatg	t gat	ttat	gtta	a					744
# 1	<210 <211 <212 <213	L> 1' 2> P		sapie	ens												
	<400 Met 1)> 6 Val	Leu	Ser	Gly 5	Ala	Leu	Cys	Phe	Arg 10	Glu	Asp	Gln	Thr	Pro 15	Leu	
75	Ile	Ala	Gly	Met 20	Cys	Ser	Leu	Pro	Met 25	Ala	Arg	Tyr	Tyr	Ile 30	Ile	Lys	
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Tyr	Ala	Asp 35	Gln	Lys	Ala	Leu	Tyr 40		Arg	Asp	Gly	Gln 45		Leu	Val	
	Gly	Asp 50		Val	Ala	Asp	Asn 55		Cys	Ala	Glu	Lys 60	Ile	Cys	Ile	Leu	
	Pro 65	Asn	Arg	Gly	Leu	Ala 70	Arg	Thr	. Lys	Val	Pro 75		Phe	Leu	Gly	Ile 80	
	Gln	Gly	Gly	Ser	Arg 85		Leu	Ala	Cys	Val 90		Thr	Glu	. Glu	Gly 95	Pro	
	Ser	Leu	Gln	Leu 100		Asp	Val	Asn	ı Ile 105		. Glu	Leu	Tyr	Lys 110		gly	
	Glu	Glu	Ala 115		· Arg	Phe	Thr	Phe 120		e Gln	. Ser	Ser	Ser 125		ser Ser	Ala	
	Phe	Arg		Glu	. Ala	. Ala	Ala 135		Pro	Gly	Trp	Phe 140		Cys	s Gly	7 Pro	
	Ala	Glu	ı Pro	Gln	Glr	Pro	Val	. Glr	ı Let	ı Thr	Lys	Glu	ı Ser	Glu	ı Pro	Ser	

Ala Arg Thr Lys Phe Tyr Phe Glu Gln Ser Trp 165 170

<210> 7

<211> 269

<212> PRT

<213> Homo sapiens

<400> 7

12 E S

l, i

iΞ

. .

Met Ala Glu Val Pro Lys Leu Ala Ser Glu Met Met Ala Tyr Tyr Ser 1 5 10 15

Gly Asn Glu Asp Asp Leu Phe Phe Glu Ala Asp Gly Pro Lys Gln Met 20 25 30

Lys Cys Ser Phe Gln Asp Leu Asp Leu Cys Pro Leu Asp Gly Gly Ile 35 40 45

Gln Leu Arg Ile Ser Asp His His Tyr Ser Lys Gly Phe Arg Gln Ala 50 55 60

Ala Ser Val Val Val Ala Met Asp Lys Leu Arg Lys Met Leu Val Pro 65 70 75 80

Cys Pro Gln Thr Phe Gln Glu Asn Asp Leu Ser Thr Phe Phe Pro Phe 85 90 95

Ile Phe Glu Glu Glu Pro Ile Phe Phe Asp Thr Trp Asp Asn Glu Ala
100 105 110

Tyr Val His Asp Ala Pro Val Arg Ser Leu Asn Cys Thr Leu Arg Asp 115 120 125

Ser Gln Gln Lys Ser Leu Val Met Ser Gly Pro Tyr Glu Leu Lys Ala 130 135 140

Leu His Leu Gln Gly Gln Asp Met Glu Gln Gln Val Val Phe Ser Met 145 150 155 160

Ser Phe Val Gln Gly Glu Glu Ser Asn Asp Lys Ile Pro Val Ala Leu 165 170 175

Gly Leu Lys Glu Lys Asn Leu Tyr Leu Ser Cys Val Leu Lys Asp Asp 180 185 190

Lys Pro Thr Leu Gln Leu Glu Ser Val Asp Pro Lys Asn Tyr Pro Lys 195 200 205

Lys Lys Met Glu Lys Arg Phe Val Phe Asn Lys Ile Glu Ile Asn Asn 210 215 220

Lys Leu Glu Phe Glu Ser Ala Gln Phe Pro Asn Trp Tyr Ile Ser Thr 225 230 235 240

Ser Gln Ala Glu Asn Met Pro Val Phe Leu Gly Gly Thr Lys Gly Gly 250 245 Gln Asp Ile Thr Asp Phe Thr Met Gln Phe Val Ser Ser 265 <210> 8 <211> 153 <212> PRT <213> Homo sapiens Ala Pro Val Arg Ser Leu Asn Cys Thr Leu Arg Asp Ser Gln Gln Lys Ser Leu Val Met Ser Gly Pro Tyr Glu Leu Lys Ala Leu His Leu Gln 25 Gly Gln Asp Met Glu Gln Gln Val Val Phe Ser Met Ser Phe Val Gln Gly Glu Glu Ser Asn Asp Lys Ile Pro Val Ala Leu Gly Leu Lys Glu Lys Asn Leu Tyr Leu Ser Cys Val Leu Lys Asp Asp Lys Pro Thr Leu Gln Leu Glu Ser Val Asp Pro Lys Asn Tyr Pro Lys Lys Lys Met Glu 90 Lys Arg Phe Val Phe Asn Lys Ile Glu Ile Asn Asn Lys Leu Glu Phe 105 Glu Ser Ala Gln Phe Pro Asn Trp Tyr Ile Ser Thr Ser Gln Ala Glu 120 Asn Met Pro Val Phe Leu Gly Gly Thr Lys Gly Gly Gln Asp Ile Thr 140 Asp Phe Thr Met Gln Phe Val Ser Ser 145 150 <210> 9 <211> 177 <212> PRT <213> Homo sapiens Met Glu Ile Cys Arg Gly Leu Arg Ser His Leu Ile Thr Leu Leu Leu Phe Leu Phe His Ser Glu Thr Ile Cys Arg Pro Ser Gly Arg Lys Ser

Ser Lys Ile Gln Ala Phe Arg Ile Trp Asp Val Asn Gln Lys Thr Phe

Tyr Leu Arg Asn Asn Gln Leu Val Ala Gly Tyr Leu Gln Gly Pro Asn 50 55 60

Val Asn Leu Glu Glu Lys Ile Asp Val Val Pro Ile Glu Pro His Ala 65 70 75 80

Leu Phe Leu Gly Ile His Gly Gly Lys Met Cys Leu Ser Cys Val Lys
85 90 95

Ser Gly Asp Glu Thr Arg Leu Gln Leu Glu Ala Val Asn Ile Thr Asp 100 105 110

Leu Ser Glu Asn Arg Lys Gln Asp Lys Arg Phe Ala Phe Ile Arg Ser 115 120 125

Asp Ser Gly Pro Thr Thr Ser Phe Glu Ser Ala Ala Cys Pro Gly Trp 130 135 140

Phe Leu Cys Thr Ala Met Glu Ala Asp Gln Pro Val Ser Leu Thr Asn 145 150 150 160

Met Pro Asp Glu Gly Val Met Val Thr Lys Phe Tyr Phe Gln Glu Asp 165 170 175

Glu

Ęij

<210> 10

-- <211> 155

<212> PRT

<213> Homo sapiens

<400> 10

Met Val Leu Ser Gly Ala Leu Cys Phe Arg Met Lys Asp Ser Ala Leu

1 5 10 15

Lys Val Leu Tyr Leu His Asn Asn Gln Leu Leu Ala Gly Gly Leu His 20 25 30

Ala Gly Lys Val Ile Lys Gly Glu Glu Ile Ser Val Val Pro Asn Arg 35 40 45

Trp Leu Asp Ala Ser Leu Ser Pro Val Ile Leu Gly Val Gln Gly Gly 50 55 60

Ser Gln Cys Leu Ser Cys Gly Val Gly Gln Glu Pro Thr Leu Thr Leu 65 70 75 80

Glu Pro Val Asn Ile Met Glu Leu Tyr Leu Gly Ala Lys Glu Ser Lys 85 90 95

Ser Phe Thr Phe Tyr Arg Arg Asp Met Gly Leu Thr Ser Ser Phe Glu 100 105 110 Ser Ala Ala Tyr Pro Gly Trp Phe Leu Cys Thr Val Pro Glu Ala Asp 120 Gln Pro Val Arg Leu Thr Gln Leu Pro Glu Asn Gly Gly Trp Asn Ala 135 Pro Ile Thr Asp Phe Tyr Phe Gln Gln Cys Asp 150 <210> 11 <211> 178 <212> PRT <213> Homo sapiens <400> 11 Met Ser Phe Val Gly Glu Asn Ser Gly Val Lys Met Gly Ser Glu Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu Glu Asp Pro Ala Gly Ser Pro Leu Glu Pro Gly Pro Ser Leu Pro Thr Met Asn Phe Val His Thr 40 35 Lys Ile Phe Phe Ala Leu Ala Ser Ser Leu Ser Ser Ala Ser Ala Glu Lys Gly Ser Pro Ile Leu Leu Gly Val Ser Lys Gly Glu Phe Cys Leu 75 70 Tyr Cys Asp Lys Asp Lys Gly Gln Ser His Pro Ser Leu Gln Leu Lys Lys Glu Lys Leu Met Lys Leu Ala Ala Gln Lys Glu Ser Ala Arg Arg 100 Pro Phe Ile Phe Tyr Arg Ala Gln Val Gly Ser Trp Asn Met Leu Glu 125 115 Ser Ala Ala His Pro Gly Trp Phe Ile Cys Thr Ser Cys Asn Cys Asn Glu Pro Val Gly Val Thr Asp Lys Phe Glu Asn Arg Lys His Ile Glu 155

Phe Ser Phe Gln Pro Val Cys Lys Ala Glu Met Ser Pro Ser Glu Val

170

Ser Asp

<210> 12 <211> 218 <212> PRT <213> Homo sapiens

165

```
15
```

Met Ser Phe Val Gly Glu Asn Ser Gly Val Lys Met Gly Ser Glu Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu Glu Asp Pro Ala Gly Ser Pro Leu Glu Pro Gly Pro Ser Leu Pro Thr Met Asn Phe Val His Thr Ser Pro Lys Val Lys Asn Leu Asn Pro Lys Lys Phe Ser Ile His Asp Gln Asp His Lys Val Leu Val Leu Asp Ser Gly Asn Leu Ile Ala Val Pro Asp Lys Asn Tyr Ile Arg Pro Glu Ile Phe Phe Ala Leu Ala Ser Ser Leu Ser Ser Ala Ser Ala Glu Lys Gly Ser Pro Ile Leu Leu Gly 105 Val Ser Lys Gly Glu Phe Cys Leu Tyr Cys Asp Lys Asp Lys Gly Gln 120 Ser His Pro Ser Leu Gln Leu Lys Lys Glu Lys Leu Met Lys Leu Ala 130 Ala Gln Lys Glu Ser Ala Arg Arg Pro Phe Ile Phe Tyr Arg Ala Gln 155 Val Gly Ser Trp Asn Met Leu Glu Ser Ala Ala His Pro Gly Trp Phe 170 Ile Cys Thr Ser Cys Asn Cys Asn Glu Pro Val Gly Val Thr Asp Lys 185 180 Phe Glu Asn Arg Lys His Ile Glu Phe Ser Phe Gln Pro Val Cys Lys 200 Ala Glu Met Ser Pro Ser Glu Val Ser Asp 215 210 <210> 13 <211> 192 <212> PRT <213> Homo sapiens <400> 13 Met Ser Gly Cys Asp Arg Arg Glu Thr Glu Thr Lys Gly Lys Asn Ser Phe Lys Lys Arg Leu Arg Gly Pro Lys Val Lys Asn Leu Asn Pro Lys

	Lys	Phe	Ser 35	Ile	His	Asp	Gln	Asp 40	His	Lys	Val	Leu	Val 45	Leu	Asp	Ser
	Gly	Asn 50	Leu	Ile	Ala	Val	Pro 55	Asp	Lys	Asn	Tyr	Ile 60	Arg	Pro	Glu	Ile
	Phe 65	Phe	Ala	Leu	Ala	Ser 70	Ser	Leu	Ser	Ser	Ala 75	Ser	Ala	Glu	Lys	Gly 80
	Ser	Pro	Ile	Leu	Leu 85	Gly	Val	Ser	Lys	Gly 90	Glu	Phe	Cys	Leu	Tyr 95	Cys
	Asp	Lys	Asp	Lys 100	Gly	Gln	Ser	His	Pro 105	Ser	Leu	Gln	Leu	Lys 110	Lys	Glu
	Lys	Leu	Met 115	Lys	Leu	Ala	Ala	Gln 120	Lys	Glu	Ser	Ala	Arg 125	Arg	Pro	Phe
	Ile	Phe 130	Tyr	Arg	Ala	Gln	Val 135	Gly	Ser	Trp	Asn	Met 140	Leu	Glu	Ser	Ala
	Ala 145	His	Pro	Gly	Trp	Phe 150	Ile	Cys	Thr	Ser	Cys 155	Asn	Cys	Asn	Glu	Pro 160
74	Val	Gly	Val	Thr	Asp 165	Lys	Phe	Glu	Asn	Arg 170	Lys	His	Ile	Glu	Phe 175	Ser
10 10 10 10 10 10 10 10 10 10 10 10 10 1	Phe	Gln	Pro	Val 180	Cys	Lys	Ala	Glu	Met 185	Ser	Pro	Ser	Glu	Val 190	Ser	Asp
7	<21	0> 1	4													
	<211> 169 <212> PRT <213> Homo sapiens															
	<40 Met		4 Gly	Thr	Pro	Gly	Asp	Ala	Asp	Gly 10		Gly	Arg	Ala	Val 15	Tyr
	Gln	Ser	Met	Cys 20		Pro	Ile	Thr	Gly 25		Ile	. Asn	Asp	Leu 30		Gln
	Gln	. Val	Trp		Leu	Gln	Gly	Glr 40		. Leu	ı Val	Ala	val 45		Arg	Ser
	Asp	Ser 50		Thr	Pro	Val	. Thr 55		. Ala	ı Val	. Ile	thr		. Lys	Tyr	Pro
	Glu 65		Leu	ı Glu	ı Gln	Gl _y		g Gl	/ Asp	Pro	75		. Lev	ı Gly	7 Ile	Gln 80

Asn Pro Glu Met Cys Leu Tyr Cys Glu Lys Val Gly Glu Gln Pro Thr 85 90 95

Leu Gln Leu Lys Glu Gln Lys Ile Met Asp Leu Tyr Gly Gln Pro Glu Pro Val Lys Pro Phe Leu Phe Tyr Arg Ala Lys Thr Gly Arg Thr Ser 125 120 Thr Leu Glu Ser Val Ala Phe Pro Asp Trp Phe Ile Ala Ser Ser Lys 135 Arg Asp Gln Pro Ile Ile Leu Thr Ser Glu Leu Gly Lys Ser Tyr Asn 155 150 Thr Ala Phe Glu Leu Asn Ile Asn Asp <210> 15 <211> 208 <212> PRT <213> Homo sapiens <400> 15 Met Arg Gly Thr Pro Gly Asp Ala Asp Gly Gly Arg Ala Val Tyr Gln Ser Ser Glu Ser Asn Ala Val Gly Met Gly Leu Trp Arg Leu Arg 20 Pro Ser Ala Leu Thr Leu Ser Pro Val Glu Ala Pro Ala Phe Ser Ala Pro Leu Cys Thr Leu Pro Phe Pro Pro Val Cys Lys Pro Ile Thr Gly 55 Thr Ile Asn Asp Leu Asn Gln Gln Val Trp Thr Leu Gln Gly Gln Asn 70 65 Leu Val Ala Val Pro Arg Ser Asp Ser Val Thr Pro Val Thr Val Ala 90 Val Ile Thr Cys Lys Tyr Pro Glu Ala Leu Glu Gln Gly Arg Gly Asp 105 100 Pro Ile Tyr Leu Gly Ile Gln Asn Pro Glu Met Cys Leu Tyr Cys Glu Lys Val Gly Glu Gln Pro Thr Leu Gln Leu Lys Glu Gln Lys Ile Met Asp Leu Tyr Gly Gln Pro Glu Pro Val Lys Pro Phe Leu Phe Tyr Arg 155 150 145

13

170

Ala Lys Thr Gly Arg Thr Ser Thr Leu Glu Ser Val Ala Phe Pro Asp

Trp Phe Ile Ala Ser Ser Lys Arg Asp Gln Pro Ile Ile Leu Thr Ser

180 185 190

Glu Leu Gly Lys Ser Tyr Asn Thr Ala Phe Glu Leu Asn Ile Asn Asp 195 200 205

<210> 16

<211> 158

<212> PRT

<213> Homo sapiens

<400> 16

1, 2

ļ, Ļ,

15

Met Glu Lys Ala Leu Lys Ile Asp Thr Pro Gln Gln Gly Ser Ile Gln
1 5 10 15

Asp Ile Asn His Arg Val Trp Val Leu Gln Asp Gln Thr Leu Ile Ala 20 25 30

Val Pro Arg Lys Asp Arg Met Ser Pro Val Thr Ile Ala Leu Ile Ser

Cys Arg His Val Glu Thr Leu Glu Lys Asp Arg Gly Asn Pro Ile Tyr 50 55 60

Leu Gly Leu Asn Gly Leu Asn Leu Cys Leu Met Cys Ala Lys Val Gly 65 70 75 80

Asp Gln Pro Thr Leu Gln Leu Lys Glu Lys Asp Ile Met Asp Leu Tyr
85 90 95

Asn Gln Pro Glu Pro Val Lys Ser Phe Leu Phe Tyr His Ser Gln Ser 100 105 110

Gly Arg Asn Ser Thr Phe Glu Ser Val Ala Phe Pro Gly Trp Phe Ile 115 120 125

Ala Val Ser Ser Glu Gly Gly Cys Pro Leu Ile Leu Thr Gln Glu Leu 130 135 140

Gly Lys Ala Asn Thr Thr Asp Phe Gly Leu Thr Met Leu Phe 145 150 155

<210> 17

<211> 157

<212> PRT

<213> Homo sapiens

<400> 17

Met Asn Pro Gln Arg Glu Ala Ala Pro Lys Ser Tyr Ala Ile Arg Asp 1 5 10 15

Ser Arg Gln Met Val Trp Val Leu Ser Gly Asn Ser Leu Ile Ala Ala 20 25 30

```
Pro Leu Ser Arg Ser Ile Lys Pro Val Thr Leu His Leu Ile Ala Cys
  Arg Asp Thr Glu Phe Ser Asp Lys Glu Lys Gly Asn Met Val Tyr Leu
                           55
  Gly Ile Lys Gly Lys Asp Leu Cys Leu Phe Cys Ala Glu Ile Gln Gly
  Lys Pro Thr Leu Gln Leu Lys Glu Lys Asn Ile Met Asp Leu Tyr Val
  Glu Lys Lys Ala Gln Lys Pro Phe Leu Phe Phe His Asn Lys Glu Gly
                                  105
  Ser Thr Ser Val Phe Gln Ser Val Ser Tyr Pro Gly Trp Phe Ile Ala
                              120
   Thr Ser Thr Thr Ser Gly Gln Pro Ile Phe Leu Thr Lys Glu Arg Gly
                          135
   Ile Thr Asn Asn Thr Asn Phe Tyr Leu Asp Ser Val Glu
                                          155
                   150
   145
<210> 18
<211> 11
< <212> PRT
<213> Human immunodeficiency virus type 1
(400> 18
   Tyr Gly Arg Lys Lys Arg Arg Gln Arg Arg
<210> 19
<211> 15
   <212> PRT
   <213> Artificial Sequence
   <223> Description of Artificial Sequence: internalizing
         domain derived from HIV tat protein
   Gly Gly Gly Tyr Gly Arg Lys Lys Arg Arg Gln Arg Arg Arg
   <210> 20
   <211> 23
   <212> DNA
   <213> Artificial Sequence
   <220>
   <223> Description of Artificial Sequence:
```

C)

Oligonucleotide 2349-98

	<400> 20 cacacgette acettette cag	23
	<210> 21 <211> 23 <212> DNA <213> Artificial Sequence	
	<220> <223> Description of Artificial Sequence: Oligonucleotide 2349-99	
	<400> 21 taaaacttgg tacgggctga ggg	23
2 H 2 H 2 H 2 H 2 H 2 H 2 H 2 H 2 H 2 H	<210> 22 <211> 23 <212> DNA <213> Artificial Sequence	
	<220> <223> Description of Artificial Sequence: Oligonucleotide 1572-36	
	<400> 22 gtgtggaatt gtgagcggat aac	23
H 18 18 18 18 18 18 18 18 18 18 18 18 18	<210> 23 <211> 23 <212> DNA <213> Artificial Sequence	
	<220> <223> Description of Artificial Sequence: Oligonucleotide 2328-91	
	<400> 23 ctatgaccat gattacgcca agc	23
	<210> 24 <211> 20 <212> DNA <213> Artificial Sequence	
	<220> <223> Description of Artificial Sequence: Oligonucleotide 2351-47	
	<400> 24 gctgtactgg ctgctggggc	20

```
<210> 25
   <211> 21
   <212> DNA
   <213> Artificial Sequence
   <220>
   <223> Description of Artificial Sequence:
         Oligonucleotide 2351-48
   <400> 25
                                                                       21
   ccttcaggct tgaggctgct g
   <210> 26
   <211> 21
   <212> DNA
   <213> Artificial Sequence
   <220>
   <223> Description of Artificial Sequence:
         Oligonucleotide 2329-93
   <400> 26
                                                                       21
   cgggcctctt cgctattacg c
r Li
::::::
<210> 27
<211> 21
| <212> DNA
   <213> Artificial Sequence
::
   <220>
   <223> Description of Artificial Sequence:
Oligonucleotide 2363-04
<400> 27
                                                                       21
📋 cctggctggt tcctgtgtgg c
   <210> 28
   <211> 21
   <212> DNA
   <213> Artificial Sequence
   <223> Description of Artificial Sequence:
         Oligonucleotide 2329-94
   <400> 28
                                                                       21
   tggcgaaagg gggatgtgct g
   <210> 29
   <211> 27
   <212> DNA
   <213> Artificial Sequence
```

	<220> <223> I	Description of Artificial S Dligonucleotide AP-1	Sequence:					
	<400> 2	29 taat acgactcact atagggc		27				
	<210> 3 <211> 3 <212> 3 <213> 3	22						
		Description of Artificial (Oligonucleotide 2353-87	Sequence:					
		<400> 30 ccttggtgag ctgtactggc tg						
	<210> <211> <212> <213>	19						
	<223>	Description of Artificial Oligonucleotide 2349-52	Sequence:					
	<400> ccgggc	31 caca caggaacca		19				
	<210><211><212><212><213>	24						
	<220> <223>	Description of Artificial Oligonucleotide 2349-51	Sequence:					
	<400> 32 aagaggccac acgcttcacc ttct							
	<210><211><212><212><213>	26						
	<220> <223>	Description of Artificial Oligonucleotide 2557-95	Sequence:					

		400> 33 agccttttt cttctttgcc tcagtg												26			
	<210> 34 <211> 26 <212> DNA <213> Artificial Sequence																
	<pre><223> Description of Artificial Sequence: Oligonucleotide 2557-96</pre>																
	<400> 34 tgccatttaa tgtaacacgg tcacag													26			
	<210> 35 <211> 459 <212> DNA <213> Mus musculus																
A THE RESIDENCE OF THE THE STATE OF THE STAT	<220> <221> CDS <222> (1)(459)																
	atg)> 35 tgc Cys	tcc	ctt Leu	ccc Pro 5	atg Met	gca Ala	aga Arg	tac Tyr	tac Tyr 10	ata Ile	atc Ile	aag Lys	gat Asp	gca Ala 15	cat His	48
	caa Gln	aag Lys	gct Ala	ttg Leu 20	tac Tyr	aca Thr	cgg Arg	aat Asn	ggc Gly 25	cag Gln	ctc Leu	ctg Leu	ctg Leu	gga Gly 30	gac Asp	cct Pro	96
	gat Asp	tca Ser	gac Asp 35	aat Asn	tat Tyr	agt Ser	cca Pro	gag Glu 40	aag Lys	gtc Val	tgt Cys	atc Ile	ctt Leu 45	cct Pro	aac Asn	cga Arg	144
	ggc Gly	cta Leu 50	gac Asp	cgc Arg	tcc Ser	aag Lys	gtc Val 55	ccc Pro	atc Ile	ttc Phe	ctg Leu	60 Gly 999	atg Met	cag Gln	gga Gly	gga Gly	192
	agt Ser 65	tgc Cys	tgc Cys	ctg Leu	gcg Ala	tgt Cys 70	gta Val	aag Lys	aca Thr	aga Arg	gag Glu 75	gga Gly	cct Pro	ctc Leu	ctg Leu	cag Gln 80	240
	ctg Leu	gag Glu	gat Asp	gtg Val	aac Asn 85	atc Ile	gag Glu	gac Asp	cta Leu	tac Tyr 90	aag Lys	gga Gly	ggt Gly	gaa Glu	caa Gln 95	acc Thr	288
		cgt Arg															336
		gct Ala															384

115 120 125

cag cag cca gtg cag ctc acc aaa gag agt gaa ccc tcc acc cat act
Gln Gln Pro Val Gln Leu Thr Lys Glu Ser Glu Pro Ser Thr His Thr
130 135 140

gaa ttc tac ttt gag atg agt cgg taa 459

Glu Phe Tyr Phe Glu Met Ser Arg 145 150

<210> 36 <211> 152 <212> PRT <213> Mus musculus

ŧΕ

Gln Lys Ala Leu Tyr Thr Arg Asn Gly Gln Leu Leu Leu Gly Asp Pro 20 25 30

Asp Ser Asp Asn Tyr Ser Pro Glu Lys Val Cys Ile Leu Pro Asn Arg 35 40 45

Gly Leu Asp Arg Ser Lys Val Pro Ile Phe Leu Gly Met Gln Gly Gly
50 55 60

Ser Cys Cys Leu Ala Cys Val Lys Thr Arg Glu Gly Pro Leu Leu Gln 65 70 75 80

Leu Glu Asp Val Asn Ile Glu Asp Leu Tyr Lys Gly Gly Glu Gln Thr 85 90 95

Thr Arg Phe Thr Phe Phe Gln Arg Ser Leu Gly Ser Ala Phe Arg Leu 100 105 110

Glu Ala Ala Cys Pro Gly Trp Phe Leu Cys Gly Pro Ala Glu Pro 115 120 125

Gln Gln Pro Val Gln Leu Thr Lys Glu Ser Glu Pro Ser Thr His Thr 130 135 140

Glu Phe Tyr Phe Glu Met Ser Arg 145 150

<210> 37 <211> 13017 <212> DNA <213> Mus musculus <220> <221> exon

<222> (8025)..(8054)

<220> <221> exon <222> (9548)..(9631) <220> <221> exon <222> (9967)..(10092) <220> <221> exon <222> (10358)..(10570) <400> 37 actaqtctcc catagacaac agctgaatgt acgaggtcag aagcaaggcc tgccccagaa 60 ccattgcaag ccaggtgctg tcttgattgt agcctcataa aaaactgatg cagaattgcc 120 ccaccaacat gctccagatt cctgctccac agaaaccctg tgaactaacc atgttgcttt 180 tagattctqc aqtaaqttqa taatctgcag taaataacat tcgatgaaag agaaacatgt 240 qtaqttactt tattatgatc aaaactttat ttctccactc tttccatttt ccttctcaga 300 attgacacca gcctttcact aacccaaata gcctatttaa atgctgatca tacttctctt 360 gttaactgtt acctgttccc aaaaggtaca attccctttc gaccatagct gcatctccca 420 cctgcacacc aggatgtttc tcatatttct acctaaaaca ttggggacta caagtgaaag 480 caaaagaggg ggtccatatc agaaccccag gtatttagct gtaaaactca cttgtcaggc 540 cagcttgaca ggtttacagt ttgtagaagg accagaaaga aggtagccaa gacagaagag 600 gcaacctctg cttgtcctag aaccttcagt ccatatacat ctaagctccc cagcaccatt 660 tctaccacag acctctcaga gttcctgagg atgcagaccc caggacactg acctcagttt 720 ccaggcaggg tttctgcaca cccccttcac actgcctgac tgggagttag tctcatggtg 780 caacactact ttgggacact gtacccatcc cctcgaccta cagaaaccat tcacttttca 840 aggtcacctc ctataggaag tatttgaaaa gatgagagtc atggtcattt gctatgataa 900 tattctgtgc ttatctccct gtaaaaagtt ggcttggggt ctctggcatg catctgacct 960 taaggttgga gctgcaccaa tatgttttta agcacccggc ataatgcttc gcaaaatttc 1020 agaacatggt ttgtacagaa tgtactttcc tccactcata caaacccttg taaaagagta 1080 gtttgaatcc caactcattc ttgaaggcca ccttttgtag ggtgacagaa tttaaaaata 1140 cagaatttaa aaatacttta tcccagggaa gctcacactt ctaaatccag aatgaaagaa 1200 gaaatagaaa cacacttgtg gtggcggtgg tggtggtgat ggtggtcgtg gtggtggtgg 1260 tggtggtggt ggtgatggtg gtggtggtgg tggtggtggt ggtcgtggtg gtgtaatgat 1320

cacagtaaag tgaggcatca tggcctgaga gagtcaggca tcacagctat tcaagtgaaa 1380 actacctact actgatttta gagttctata attttagtag cagccacagg cctggggcct 1440 gggcctatat tttcagagag gaaatgttca cagcaggtca actgcagaca gtgaagatca 1500 gaaatgtttc ataatcaggt catcagagaa aaggcaaagg agctgatgga ctttatcctg 1560 aaaaagcaaa atccaaccca cctcatgctt aatgcattca aaggtctgcg ggcagaagaa 1620 tacattttgc tttttattat tataaattac ctggagaata tttttgtctg aattatctcc 1680 caaatattaa ccataaaaat aaaaaattcc atgtgtgctt ctcccagggg ctataaagcc 1740 cctggtctta gagttgttgg ggcaaaacct gacctttgaa gtagttactt ttgaagatgc 1800 cataccatac atttggccac ttggagagag tctaatgtca catctaaagg gttactctga 1860 tgctctgttt tctcatatgc ccttggctta cagctaacta tggctccagc taaactataa 1920 agttccttgg caacagagat ggtacgctat gtgtctttga cacagcagaa taaatgctta 1980 gtgaacatta ctgattgcct gacaggacac ctcacacttt ggtactttca acagagggat 2040 gtaaacttat gaagaacaat gaagaatgaa tattggcaat aaaagcaaaa attggttaac 2100 ccaattctag ctctgaaatc atttttaggt agtgggaagt ctttttgttt tgtttattca 2160 ctttacatcc caattgctgt cctccctcca agttccccac caccaccaca gtcctttttc 2220 cctccccttc tcctctgaga gaatggagaa ccctcctgga tattccccca tcatgaaaca 2280 ttaagtetet geagggetag acaetteece cagtgaggee agteagggea geecagetag 2340 aaaaagcata teecacagae agacaacage ttttgggata geecegttee agttgtttag 2400 gatccacatg aaggctgagc tgcacatctg ctacatatga atgaggaggc ctaggtccag 2460 cctgtgtatg ttctttggtt ggtggttcag actctgagag ccccaagggt ccaggtcagt 2520 tgactctgtt ggtcttcctg tggacaccct gtccccttcc agcccacaat ccttccccta 2580 atccttctcc ttctcacttc cataagagtg tgaggagtct ttaaaaacat gaagcatttt 2640 atctccccag ggcaacacat ggaaatgaaa gattgtgaaa agtaatttaa agaaaaagaa 2700 aaaaaaattt aacaaggaat aagaatcttg tttctctgaa aatgcttaag agtgtggaaa 2760 acataaactg gattctaata gaatgcaatt ggattgtaat gaaaacctat caaagttatg 2820 aaatagcttt cactaccttg cacaaaatct cttggcatgt gtgttgttgg caaattttct 2880 tgttagttta aaaccacaac aataacaaca aaatagcaaa aattgggtct cagcctcatt 2940 cattttttct catttcttgc tctgtgatcg tctgggtctt aagctgacac ctcaccaatt 3000 cctcatcaag acctttgtgg aaatttgcaa atgtcccaaa aaggagaatt acaataagtc 3060 aqaqaacqtt ctgtccaatt ctttatccct agtgatggat gagtaaagga tgtataagag 3120 atggataaat ggactgatgt acagataaat gaaggaatat gtacatggtt aggtggatag 3180 atgacttact caacagatga gtagaaggat gagaaataga tggacagctg gactgaggca 3240 tgcaaagtca actggagaac tgagtctctt gaccatgcac tgtccagggt ctcatattcc 3300 ctagagtcca gggcccatgg ctcctgtgcc atccccatgc aaatctaagg ttaatacgtt 3360 ctacagctga gtttccttac atatgtgtct cagtaagttt gtatcaacta attaaatctg 3420 aaaggagttc cttctgatct tcccaaacag agccacactc gtgatgaagt cagccctgct 3480 tcattgtggt tctctggatg catctggctt ccatcagcat aatctttcta ttcttgatcc 3540 ttccaacctc ttcaggtctc agacagaacc ccatggagca tcaaagaggt ttgaccccag 3600 cattgtttat gtagctgcaa aaccactaat aacacagtca atgacagtag ctacagagac 3660 agcaggtcag tgtctggcct ctgtcaaggc tttatgagtg actctctccc cttcccgcaa 3720 atactcatta atctccccac ctccttatta tttggactgt gttgaagata ttatgaaatc 3780 tctgggctct tcttcccgga tctagagcca attacagatt ctgtaggttt gacccaccct 3840 gaccagacat tataaacaca gtgctggtgc cctgaagaaa acagttggag actccaggca 3900 ttagaatcca ggcaccagga actacaggtc agtggtgaca gtcggtctct ctgtgtatct 3960 cttacacaca cacacataca cacacacaac acaacataca cacacataca acacacaaca 4020 catacacata caacacatac acacacaca cacttttctg taatgtctcc aaaattctca 4080 ggctctaggg aagaagaaat gtcttttaga gaatgcggtg tgatgttcta taagtctagg 4140 aatacttgat agaatttaat gagaagtata gattaggtca aagcaagggt actacatatt 4200 tggaaccaca gagttttgaa agtcatctca aaagaaatta tttaggccag agatgttcaa 4260 aaaatgtttt gtttgtgaca tatggaagct cccatggaga cattctgtga ttctcatcaa 4320 tagacagtag ggatgccacc aaggtgctaa cgtcttcatc accccatcat ctatcataca 4380 tccaaatggt ttctttgaaa acaatctcct tgtgaaactt aaagtagcct tgaaaatata 4440 ataatcttgt ccagcctctc atttcaatgg gaatagattg aaggcctaag gaccaaaaca 4500 taagttattt ttagaatcca gcctttcagt caaagcttga ttcatgcata tctgtgttct 4620 gatcttaagg tgctgtgtct gtcagttgta tagttggata gaggtacaga tgagctatat 4680 acatcatgct tcaagatttc aggatcttat aacttttata aagcaaataa tttgtcttaa 4740

tgcacactaa taaacaatat agcaaagttt gacaggagtt cagagtactg ttagagaagt 4800 gaagggaaga attttgttat gatagtaaag gggaaaatca aattttgagt catggaatca 4860 tacatagttt gacatagaaa gaaccttggc aaccacataa tctaatgcat gagcccaaga 4920 actggcctgt gtttttaaga tctcattctc agctgttatg taactgaaca gacaagatac 4980 taagcccaag tatagtgaag ccatgtccag tgatcttaat aggagtgaca ggaatggttg 5040 gtgatgaaga ggggtggatt ttgagcagga ataccaaaag caatgctgac tgtgcccttg 5100 gagagaatta gcatgagtcc ttgagagaaa aatgagatgc tattgcacaa gcaacctagg 5160 gccagatggt gtcaagatag gtggccatcg tggactttag aaccaggcag gaatgtgatc 5220 agagatgtac tttatgtagg ttaggtttga ttcagaaacc aggagggtta gcatgtttac 5280 aatggtgact aaaaacaagc acaaggttat actttaaaga aataatctct gaaaagaagg 5340 gaggtatatt ttcagtgccg gaaagaggaa tattacaaaa gtgagaggag tagatttgag 5400 aaagagaagt ggattgtgga ggagcagatg ctcaccacgc ccttacactc acttgaactg 5460 acacccaaag atgaaggtgt gctgtggact gctgaagctc agcctgtggc tgggaagcag 5520 taaacaaaat tgctcatcac agctgtacaa gatattccat agcatataaa aataaaagtg 5580 cttaggctat tctcttacaa ctctcagcct tatgaatgac ccggaaggaa aagaactcta 5640 caatgtgcct gtgtctgttc ttacttcctc tgccacaagc aaaagagcct tgggaattgg 5700 ctcagaggga acgtcatcaa acaggctggc cttgaggctg ggctgttatt cgtctacctg 5760 ggatagagga attcgctatt cttttataat ccaagtgtgg cctggggacc agcagcatta 5820 ttaagacctg gttgcatgtt tgaaatgcag tctcagattt catcccagac ctaaagagta 5880 acactgtttt catgaggata caagattaag aaatatgcat tagagagtaa ttggctaaat 5940 gggtaaatgt catgcaagca ggaggatctg attgactccc caggacccac acagttccca 6000 tgccgtagag cacatctgta atcacagtag gcgtatgatg aaatgggagg tgaatcaaga 6060 gaatctctag cagctacggg ctggccagcc tcccatgcac agcactaaat aaggcaagga 6120 ccaatacctg aagttgtccc attaccttca catatacacc acggcatgtg tgtacttgta 6180 ctcacacata caaacaaata cacacqtgca cacatacaaa actcagagat taaggacaat 6240 tggcctgaca tatcagttcc taagcctggc tcattgcttg taacactaca agcagtatta 6300 aataaggata ggcgagagaa cagttaccga atggttcaga agtggggcca tgcctgtgac 6360 tttaaacaaa tgtttcatat ttttaaataa taacacttag attacaaaat aaatttacta 6420 caggaaaatg ttaagaacta tcaacaacca ttgactatcc tgtcggccac aaatgagtgt 6480

tataacaagc accagccgtc cttgtccaca tgtgtgtgtg tctacacagc tatgaattta 6540 attgggataa taatgtgcac attctttacg gcctgcagtt tttacttcat gtatttgaaa 6600 tgtttgtgcc acaaatgtca tctttaagga gcatatcctt atttcctgga tttatcattc 6660 cctttcagcc gactggacat tgacagcatt tccaactttt caaccttgta aaaataacta 6720 attgaactat tttataacta agcatttggg caatcaatta cctctgcctg gaatgggggc 6780 aacaacacat gcaatcatgg gaaagccagg atgctgctgt ctgatcccta gccctggcat 6840 tcqtqcaqaa cctcactctc atctgtgccc tgatatcctt cactctcaag tcttttccca 6900 gtgactttta aaggcaacag aatcatatag ccaataatga aagctacttg gtctacagtt 6960 gtgtggcgtt ttttatagat attttcttca tttacatttc aaatgctatc ccaaaagtcc 7020 cctataccct cccccaccct gctcccctac ccactcactc ccacttcttg gccctggctt 7080 tcccccttac tggggcatat aaagtttgct agaccaaggg gcctctcttc ccaatgatgg 7140 ccaactaggc cattttctgc tacatatgca gctagagaca ccagttctgg ggttactggt 7200 tagttcatat tgttgttcta cctatggggt tgcagacccc ttcagctctt gagtactttc 7260 tctagctcct ccattgggag ccctgtgttc catcctatag atgactgtga gcatccactt 7320 ctgtatttgc caggcactgg catatgaaat agtatctgca tttggtggct gattatggga 7380 tggaccccg ggtggggcag tctctggatg gtccatcctt tcatcttagc tccaaacttt 7440 gtctctgcaa cttcttccat ggatatttta gtccctaatc tagggagaaa tgaagtatcc 7500 acaagttgat cttccttctt gattttctta tgttttagaa gttgtatctt ggatattcta 7560 ggtttctggg ctaatatcca cttatcagtg agtacatatc aagtgaattc ttttgtgatt 7620 aggttacctc actcaagatg atattctcca ctatgttcat agcagcccta tttatagtag 7680 ccagaagetg gaaagaaccc agteeetcaa cagaggaatg gatacagaaa atgtggcaca 7740 tttatgcaat ggagtaccac tcagatatta aaaacaacga atttatgaaa ttctcgggca 7800 aaaccctatc taaagaccag gaataaggaa aagatggact gcctgcctgc agctgggaga 7860 gctggggaga cctttgtgga ttctgtaata cttaggggta cggaacagct tgtggctgga 7920 taattctgag ctccagcatg tctgccccc aaaaaacatt ctgtttttct gaaagccttt 7980 ttettetttg ceteagtgaa gaccagacae teccaactge agga atg tge tee ett ccc atg gca aga tac tac atgtaagtaa tcttaacgat cgctcaatca 8084 aggggcctgg agatcacatg agaagggaaa aggctgagtc aaagggacaa agctccctct 8144

agccacagaa atctcaaaca ctgaataatt gatcttcatc tttgtcaatc acaacagccc 8204 tctttcctgg tgacagaatg gaacaactgt aagagtggta ttgcttagtc cattttacag 8264 acceggaaac teaaceteca egaggttata caatttteet catgteatge aattaceeaa 8324 aagcagagag tgggatcgga ctctctgttc tctaaactga tgtagctagt tcttagaaag 8384 ctcaaacaat cttgagtccc aaggacagca cctttatggt cacctggatt gatacctata 8444 tcaaaaaaaa aaaaaggtct cactagatag ccctggctac cctgaaactc tcactgtgta 8504 catttaggtg accacgaact cacagagatc tgccttccaa gtgctgggat taaagtatgt 8564 accaccacac ctgcatcttt gacaataact gagtggtatc taaattcttc cagtggctaa 8624 acagttaagt cccagttccc aaagtctgag aaaaatgcca ggtggtgaaa tctgtacaga 8684 cctttgttct taatgtacaa gtgagcctgc tttaaaaaaca atacgcaagc tgtttttgct 8744 attgctaagt gttgcagaga cagaaaaggc tcccagaagt ggtaactttg gtccagaggt 8804 tctgttctca aactcattgt gagctctgaa agcaactgat gggcagctct gaaatcagct 8864 gggcaattag gctaataaca ggcataattt taatgtttca cacgcatgac agttcctccc 8924 cagctgccct agtacatact taccctccta ggcacgtcat tagacccata ggtataacca 8984 gtgactaatc aggccctggt ctaattctaa gttggcctcc tatataagtg ccactcagag 9044 tgtacctcat catggctgta gtgggcccag agtctaggga catagacttt tctattgtcc 9104 atcaagctca gaaacatcat ggatagggtt cattgtatct ccagggtacc tgagcttcaa 9284 agcaactcct cagacagcca tgaaaacatc ctcaattacc tcatgagaag acactattgt 9344 catttctgga gcctctgata atcctgagcc taggcagctt tgggatgaaa caatttctac 9404 ccttattgga acagtgtccc tctcctgtct ggaaacaatt caccaaaggc tccatgtggt 9464 tgtccagtaa ggtggtatgg ggacagaaat ggacaatgat ccctgagggc agtgatccat 9524 taaccttgcc ctcctatttc aga atc aag gat gca cat caa aag gct ttg tac 9577 aca cgg aat ggc cag ctc ctg ctg gga gac cct gat tca gac aat tat 9625 9681 aqt cca gqtgatcttc cggtggtggg ggtgggggag tggaggggag ggtgtggggg gggctctctt ccagaagttg cttagtgtcc atctgccaca aggccttgat tctttccttc 9741 aattgtgtct ctagagacat gagaatattg tcacagtgat aaggagaaga ggtagggca 9801 gtttcttcct gtaaaaaatg aattccattt accetgeagt etccataeag aaacaggeca 9861

gaggggggca gacccagtaa cttctagctg agccctacct tgcttaaaac ctgccatctg 9921 tggtcccctc actgtctgaa ttgcattctg tcttacctcc cagag aag gtc tgt atc 9978 ctt cct aac cga ggc cta gac cgc tcc aag gtc ccc atc ttc ctg ggg 10026 atg cag gga gga agt tgc tgc ctg gcg tgt gta aag aca aga gag gga 10074 cct ctc ctg cag ctg gag gtgagacacc cctcctcatt gcagtcagta 10122 ctgccactgg aacatagtga catctttgaa cccacatgtc ccctctcttg tttcccatct 10182 atctctcttt gcctccagct gagggactct agcctttggg gatgtacaga aagaacatgg 10242 cttcggaaaa ctcttcccta ttgagtcctt ctttggccaa gcctctgagg cactaagggc 10302 tgacgtccca accaaacact catttcatct cacagctgtc tccctttccc cacag gat 10360 gtg aac atc gag gac cta tac aag gga ggt gaa caa acc acc cgt ttc 10408 acc ttt ttc cag aga agc ttg gga tct gcc ttc agg ctt gag gct gct 10456 gcc tgc cct ggc tgg ttt ctc tgt ggc cca gct gag ccc cag cag cca 10504 gtg cag ctc acc aaa gag agt gaa ccc tcc acc cat act gaa ttc tac 10552 10600 ttt gag atg agt cgg taa agagacataa ggctggggcc tcgtctagtg cccccagtct gagatcttct tgctcagcat ctctggaaag cagaataagg aagataccaa 10660 agatgtttgg gtcttaatcc ccagaatctg tgaccgtgtt acattaaatg gcaaagggat 10720 ttttttttc cttcatggtc catttgggcc cattggaatc atctgaggcc tcatgaggag 10780 aaggaagagg tcagagggag actggggcaa actttggtac taaaagtaac aatggagaca 10840 gggaccataa gctgatgggt aacagtggtt tctagaaacc ggaaatgatg agagctctcc 10900 tgacacaggt tctggatttt tctggactga agaatggtga aataatacag ctccattatt 10960 ttaagccact gagtttgaga tcattcaatg aagctgtcat aataaaacct gtgcttcaca 11020 tacaattcaa tattggtagg caccccggtg atttcttgga aagacatcta gggattctcc 11080 tggatgctga ttccagggtc cagtggagtc cctgggttga agagatttca caacccagaa 11140 catcaggete gaetetteta aaagteegte gttgcacece ttgeetgaga geattageaa 11200 tttctatttc ataggaaatc tgtgtccctg cccctgctaa agcagggagc ctggaccgtc 11260 ctgatttagt gaggggtgag ctgctggcac ttttttgtgt caccagtgtc ttaagcagtg 11320 atggagcaca aaagatcttt actgagaaga tggccatgaa gctctggcta gacaccaaga 11380 atatgatata agcagagcta cagcacaaga tgagccaatg aggaaagcca ttcagggagg 11440 ctaagcccag cttcccaaag ggacagctaa ccctggactc aaatgaatag gggttttcct 11500 ggcagagaac ataggtcaag cattctaggt agaatcagca attcagaaag gtgtgagaga 11560 ggcatggaga gctccaggca tgtctgggct atggtgtgtc attcttgtgg caagaatcca 11620 acgtctgtgg ttaaggagtt gctgaaaatt aaaataggaa aatgggtaga gtctaattgt 11680 gaatgacttg caaaggagtt tagcccataa gtggggagct cagaggagtc atctaaggat 11740 tgcaagcagg ggccctgtga tcattgctgg accagcctag gtgctacaga gcctaccttc 11800 agetetgeat ceteacteae atceaggtae etteagaggt caatttetgt getetggtte 11860 tatgggtagc ctgaccctgt ttcatcttct tgtataactt aggcacataa gcttagggac 11920 tggtagagtt tacttgagtg attggtgaat caggcagcac caaactacaa gttgttcagg 11980 gctttaccaa gggggcactg attggagaat tggaatgagg gtggttagaa tgcattcaga 12040 aaacaagggg aagaaaaatt tgattgctta aagtggaaag tcccaactta aatgttagtc 12100 agtagtttct aattacttga gtctctaatt agaggttagt tggcagtttc tggttagtta 12160 atctaagttt cattttctta ggctatgacc attctctgag tcgcatgtta gcaatgcagt 12220 aagaactcaa gacccagaat agcctctgtt aattatttta gcaatgatca ctcatttctg 12280 ttgcctccta ttgagatctg ttcccatgga ccacccaggc acatcaggcc tcctagtacc 12340 aacataataa tgattgctgc acagacaaaa tattttttt cagtatctgg tatttgctac 12400 atttccatta gtgctggagg gaaggctaca acgaccatga aggcatggcc cctgccttct 12460 aaggacttac aatgtaatag gagccctgac attataaagt gggtcacctt gtttcaaact 12520 gagccaaact gaggctgagg gcttagatta gtggtaggtc actttccaga catgttcagt 12580 gctaagaaaa acacattctg gggttagtta gatgttttag ttcatttgat aagaagccca 12640 atgattggac tttcaacttc tggaacccat gtggtggaag agagaaccaa cttctgacca 12700 tttgggtcat ggcacatccc ctaccatcac aagaactcac caaaataaat tagaaaaatc 12760 aagaaaaact catatcctat agacctctgg tagaattagc agaacgctgc tgtggcactt 12820 gggatttgaa actcaaaaat ggaagaagct acttgtgacc gttcaagact ccagggaggc 12880 tcctctgaca catcccacga ctcaggctta aattccttct tctccctaga aggccacgcc 12940 atcttctcaa ccaggccaca gatgctataa ttatgtaaat gtgtgggaga ggcacacttt 13000 13017 agatcttatc cactagt